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TERMINAL (ENTER 1, 2, 3, OR ?):2

***** Welcome to STN International *****

NEWS 1 Web Page for STN Seminar Schedule - N. America
 NEWS 2 JAN 08 CHEMLIST enhanced with New Zealand Inventory of Chemicals
 NEWS 3 JAN 16 CA/Caplus Company Name Thesaurus enhanced and reloaded
 NEWS 4 JAN 16 IPC version 2007.01 thesaurus available on STN
 NEWS 5 JAN 16 WIIDS/WPINDEX/WPIX enhanced with IPC 8 reclassification data
 NEWS 6 JAN 22 CA/Caplus updated with revised CAS roles
 NEWS 7 JAN 22 CA/Caplus enhanced with patent applications from India
 NEWS 8 JAN 22 PHARM reloads with new search and display fields
 NEWS 9 JAN 29 CAS Registry Number crossover limit increased to 300,000 in multiple databases
 NEWS 10 FEB 15 PATDPAСП enhanced with Drug Approval numbers
 NEWS 11 FEB 15 RUSSIAPAT enhanced with pre-1994 records
 NEWS 12 FEB 23 KOREAPAT enhanced with IPC features and functionality
 NEWS 13 FEB 26 MEDLINE reloaded with enhancements
 NEWS 14 FEB 26 EMBASE enhanced with Clinical Trial Number field
 NEWS 15 FEB 26 TOXCENTER enhanced with reloaded MEDLINE
 NEWS 16 FEB 26 WIIDS/WPIX enhanced with new FRAGITSTR display format
 NEWS 17 FEB 26 CAS Registry Number crossover limit increased from 10,000 to 300,000 in multiple databases
 NEWS 18 MAR 15 WIIDS/WPIX enhanced with new FRAGITSTR display format
 NEWS 19 MAR 16 CASREACT coverage extended
 NEWS 20 MAR 20 MARPAT now updated daily
 NEWS 21 MAR 22 LWP1 reloaded
 NEWS 22 MAR 30 RDISCLOSURE reloaded with enhancements
 NEWS 23 APR 02 JICST-EPLUS removed from database clusters and STN
 NEWS 24 APR 30 GENBANK reloaded and enhanced with Genome Project ID field
 NEWS 25 APR 30 CHEMCATS enhanced with 1.2 million new records
 NEWS 26 APR 30 CA/Caplus enhanced with 1870-1889 U.S. patent records
 NEWS 27 APR 30 INPADOC replaced by INPADOCDB on STN
 NEWS 28 MAY 01 New CAS web site launched
 NEWS 29 MAY 08 CA/Caplus Indian patent publication number format defined
 NEWS 30 MAY 14 RDISCLOSURE on STN Easy enhanced with new search and display fields
 NEWS 31 MAY 21 BIOBIS reloaded and enhanced with archival data
 NEWS 32 MAY 21 TOXCENTER enhanced with BIOBIS reload
 NEWS 33 MAY 21 CA/Caplus enhanced with additional kind codes for German patents
 NEWS 34 MAY 22 CA/Caplus enhanced with IPC reclassification in Japanese patents

NEWS EXPRESS NOVEMBER 10 CURRENT WINDOWS VERSION IS V8.0.1C, CURRENT MACINTOSH VERSION IS V6.0.C(EU) AND V6.0.JC(JP), AND CURRENT DISCOVER FILE IS DATED 25 SEPTEMBER 2006.

NEWS HOURS STN Operating Hours Plus Help Desk Availability
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***** STN Columbus *****

FILE 'HOME' ENTERED AT 09:24:16 ON 23 MAY 2007

>> file reg	COST IN U.S. DOLLARS	SINCE FILE ENTRY	TOTAL SESSION
FULL ESTIMATED COST		0.21	0.21

FILE 'REGISTRY' ENTERED AT 09:24:23 ON 23 MAY 2007
 USE IS SUBJECT TO THE TERMS OF YOUR STN CUSTOMER AGREEMENT.
 PLEASE SEE "HELP USAGETERMS" FOR DETAILS.
 COPYRIGHT (C) 2007 American Chemical Society (ACS)

Property values tagged with IC are from the ZIC/VINCI data file provided by Infomach.

STRUCTURE FILE UPDATES: 22 MAY 2007 HIGHEST RN 935655-41-7
 DICTIONARY FILE UPDATES: 22 MAY 2007 HIGHEST RN 935655-41-7

New CAS Information Use Policies, enter HELP USAGETERMS for details.

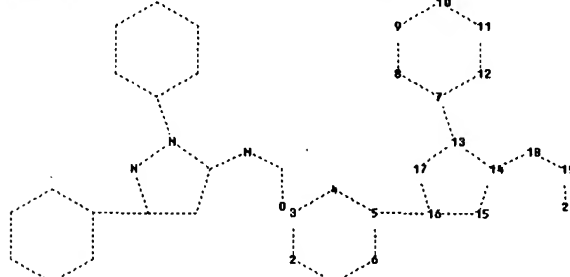
TSCA INFORMATION NOW CURRENT THROUGH December 2, 2006

Please note that search-term pricing does apply when conducting SmartSELECT searches.

REGISTRY includes numerically searchable data for experimental and predicted properties as well as tags indicating availability of experimental property data in the original document. For information on property searching in REGISTRY, refer to:

<http://www.cas.org/support/stninfo/stndoc/properties.html>

>>
 Uploading C:\Program Files\Stnxp\Queries\10.572772\no bonds.str

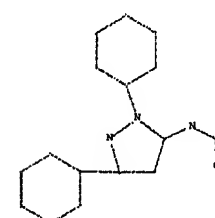


chain nodes :
 18 19 20
 ring nodes :
 1 2 3 4 5 6 7 8 9 10 11 12 13 14 15 16 17
 chain bonds :
 5-16 7-13 14-18 18-19 19-20
 ring bonds :
 1-2 1-6 2-3 3-4 4-5 5-6 5-16 7-8 7-12 8-9 9-10 10-11 11-12 13-14 13-17 14-15
 15-16 16-17
 exact/norm bonds :
 1-2 1-6 2-3 3-4 4-5 5-6 5-16 7-8 7-12 7-13 8-9 9-10 10-11 11-12 13-14
 13-17 14-15 14-18 15-16 16-17 18-19 19-20

Match level :
 1:Atom 2:Atom 3:Atom 4:Atom 5:Atom 6:Atom 7:Atom 8:Atom 9:Atom 10:Atom
 11:Atom 12:Atom 13:Atom 14:Atom 15:Atom 16:Atom 17:Atom 18:CLASS 19:CLASS 20:CLASS

L1 STRUCTURE UPLOADED

>> d
 L1 HAS NO ANSWERS
 L1 STR



Structure attributes must be viewed using STN Express query preparation.

>> s 11 SSS SAM
 SAMPLE SEARCH INITIATED 09:24:56 FILE 'REGISTRY'
 SAMPLE SCREEN SEARCH COMPLETED - 91 TO ITERATE
 100% PROCESSED 91 ITERATIONS 24 ANSWERS
 SEARCH TIME: 00.00.01
 FULL FILE PROJECTIONS: ONLINE **COMPLETE**
 BATCH **COMPLETE**
 PROJECTED ITERATIONS: 1248 TO 2392
 PROJECTED ANSWERS: 187 TO 773

L2 24 SEA SSS SAM L1

>> d scab
 'SCAB' IS NOT A VALID FORMAT FOR FILE 'REGISTRY'

The following are valid formats:

Substance information can be displayed by requesting individual fields or predefined formats. The predefined substance formats are: (RN = CAS Registry Number)

REG	- RN
SAM	- Index Name, MF, and structure - no RN
FIDE	- All substance data, except sequence data
IDE	- FIDE, but only 50 names
SQIDE	- IDE, plus sequence data
SQIDE3	- Same as SQIDE, but 3-letter amino acid codes are used
SQD	- Protein sequence data, includes RN
SQD1	- Same as SQD, but 3-letter amino acid codes are used
SQN	- Protein sequence name information, includes RN

CALC	- Table of calculated properties
EPROP	- Table of experimental properties
PROP	- EPROP and CALC

Any CA File format may be combined with any substance format to obtain CA references citing the substance. The substance formats

must be cited first. The CA File predefined formats are:

ABS -- Abstract
APPS -- Application and Priority Information
BIB -- CA Accession Number, plus Bibliographic Data
CAN -- CA Accession Number
CBIB -- CA Accession Number, plus Bibliographic Data (compressed)
IND -- Index Data
IPC -- International Patent Classification
PAT5 -- PI, SO
STD -- BIB, IPC, and NCL

IABS -- ABS, indented, with text labels
IBIB -- BIB, indented, with text labels
ISTD -- STD format, indented

OBIB ----- AN, plus Bibliographic Data (original)
OIBIB ----- OBIB, indented with text labels

SBIB ----- BIB, no citations

SIBIB ----- IBIB, no citations

The ALL format gives FIDE BIB ABS IND RE, plus sequence data when it is available.

The MAX format is the same as ALL.

The IALL format is the same as ALL with BIB ABS and IND indented, with text labels.

For additional information, please consult the following help messages:

HELP DFIELDS -- To see a complete list of individual display fields.
HELP FORMATS -- To see detailed descriptions of the predefined formats.
ENTER DISPLAY FORMAT (IDE):d scan
'D' IS NOT A VALID FORMAT FOR FILE 'REGISTRY'

The following are valid formats:

Substance information can be displayed by requesting individual fields or predefined formats. The predefined substance formats are: (RN = CAS Registry Number)

REG - RN
SAM - Index Name, MF, and structure - no RN
FIDE - All substance data, except sequence data
IDE - FIDE, but only 50 names
SQIDE - IDE, plus sequence data
SQIDE3 - Same as SQIDE, but 3-letter amino acid codes are used
SOD - Protein sequence data, includes RN
SOD3 - Same as SOD, but 3-letter amino acid codes are used
SON - Protein sequence name information, includes RN

CALC - Table of calculated properties
EPROP - Table of experimental properties
PROP - EPROP and CALC

Any CA File format may be combined with any substance format to obtain CA references citing the substance. The substance formats must be cited first. The CA File predefined formats are:

ABS -- Abstract
APPS -- Application and Priority Information
BIB -- CA Accession Number, plus Bibliographic Data

CAN -- CA Accession Number
CBIB -- CA Accession Number, plus Bibliographic Data (compressed)

IND -- Index Data
IPC -- International Patent Classification

PAT5 -- PI, SO

STD -- BIB, IPC, and NCL

IABS -- ABS, indented, with text labels
IDIB -- BIB, indented, with text labels
ISTD -- STD format, indented

OBIB ----- AN, plus Bibliographic Data (original)
OIBIB ----- OBIB, indented with text labels

SBIB ----- BIB, no citations

SIBIB ----- IBIB, no citations

The ALL format gives FIDE BIB ABS IND RE, plus sequence data when it is available.

The MAX format is the same as ALL.

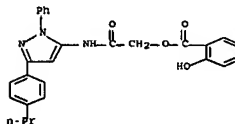
The IALL format is the same as ALL with BIB ABS and IND indented, with text labels.

For additional information, please consult the following help messages:

HELP DFIELDS -- To see a complete list of individual display fields.
HELP FORMATS -- To see detailed descriptions of the predefined formats.
ENTER DISPLAY FORMAT (IDE):end

>> d scan

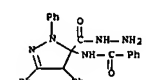
L2 24 ANSWERS REGISTRY COPYRIGHT 2007 ACS on STN
IN Benzolic acid, 2-hydroxy-, 2-oxo-2-[(1-phenyl-3-(4-propylphenyl)-1H-pyrazol-5-yl)amino]ethyl ester (9CI)
MF C27 H25 N3 O4



PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

HOW MANY MORE ANSWERS DO YOU WISH TO SCAN? (1):1

L2 24 ANSWERS REGISTRY COPYRIGHT 2007 ACS on STN
IN 1H-Pyrazole-5-carboxylic acid, 5-(benzoylamino)-4,5-dihydro-1,3,4-triphenyl-, hydrazide (9CI)
MF C29 H25 N5 O2

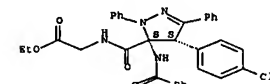


PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

HOW MANY MORE ANSWERS DO YOU WISH TO SCAN? (1):1

L2 24 ANSWERS REGISTRY COPYRIGHT 2007 ACS on STN
IN Glycine, N-[5-(benzoylamino)-4-(4-chlorophenyl)-4,5-dihydro-1,3-diphenyl-1H-pyrazol-5-yl]carbonyl-, ethyl ester, cis- (9CI)
MF C33 H29 Cl N4 O4

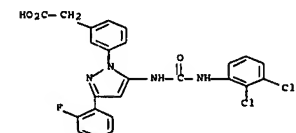
Relative stereochemistry.



PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

HOW MANY MORE ANSWERS DO YOU WISH TO SCAN? (1):1

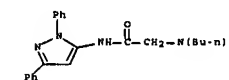
L2 24 ANSWERS REGISTRY COPYRIGHT 2007 ACS on STN
IN Benzeacetic acid, 3-[5-[(2,3-dichlorophenyl)amino]carbonyl]amino]-3-(2-fluorophenyl)-1H-pyrazol-1-yl- (9CI)
MF C24 H17 Cl2 F N4 O3



PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

HOW MANY MORE ANSWERS DO YOU WISH TO SCAN? (1):1

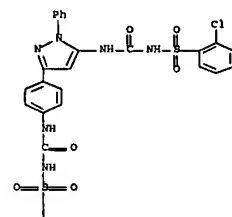
L2 24 ANSWERS REGISTRY COPYRIGHT 2007 ACS on STN
IN Acetamide, 2-(dibutylamino)-N-(1,3-diphenylpyrazol-3-yl)- (8CI)
MF C25 H32 N4 O
CI COM



PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

HOW MANY MORE ANSWERS DO YOU WISH TO SCAN? (1):1

L2 24 ANSWERS REGISTRY COPYRIGHT 2007 ACS on STN
IN Benzenesulfonamide, 2-chloro-N-[(3-[(4-[(2-chlorophenyl)sulfonyl]amino)carbonyl]amino)phenyl]-1-phenyl-1H-pyrazol-5-yl]amino]carbonyl- (9CI)
MF C29 H22 Cl2 N6 O6 S2



PAGE 1-A



PAGE 2-A

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

HOW MANY MORE ANSWERS DO YOU WISH TO SCAN? (1):0

-> s 11 sess full
FULL SEARCH INITIATED 09:25:41 FILE 'REGISTRY'
FULL SCREEN SEARCH COMPLETED - 1599 TO ITERATE

100.0% PROCESSED 1599 ITERATIONS 413 ANSWERS
SEARCH TIME: 00:00.01

L3 413 SEA SSS FUL L1
> file hcplus COST IN U.S. DOLLARS SINCE FILE TOTAL
ENTRY SESSION
FULL ESTIMATED COST 172.55 172.76

FILE 'HCPLUS' ENTERED AT 09:25:46 ON 23 MAY 2007
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FILE COVERD 1907 - 23 May 2007 VOL 146 ISS 22
FILE LAST UPDATED: 22 May 2007 (20070522/ED)

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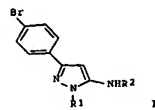
This file contains CAS Registry Numbers for easy and accurate substance identification.

-> s 13
L4 128 L3
>> s 14 and py <2004
23932542 PY <2004
L5 92 L4 AND PY <2004

-> d bib abr hitstr 1-10

L5 ANSWER 1 OF 92 HCPLUS COPYRIGHT 2007 ACS on STN
ACCESSION NUMBER: 2004:153241 HCPLUS Full-text
DOCUMENT NUMBER: 141:395467
TITLE: 3-(p-Bromophenyl)-5-aminopyrazole and some derivatives
AUTHOR(S): Nam, N. L.; Grandberg, I. I.; Sorokin, V. I.
CORPORATE SOURCE: Kafedra Org. Khim., Timiryazevsk. S.-Kh. Akad., Russia
SOURCE: Izvestiya Timiryazevskoi Sel'skogozyaistvennoi

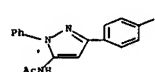
Akademii (2003), (4), 142-146
CODEN: ITSAAT; ISSN: 0021-342X
DOCUMENT TYPE: ANO "Izdatel'stvo MSKhA"
LANGUAGE: Journal
OTHER SOURCE (S): Russian
GI CASREACT 141:395467



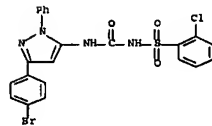
AB 5-Aminopyrazoles I (R1 = H, Me, Ph; R2 = H) were readily prepared via cyanation of α , β -dibromoacetophenone with sodium cyanide followed by heterocyclization of 4-bromo- α -cyanacetophenone with the corresponding hydrazines. Pyrazole I (R1 = Ph; R2 = H) was further functionalized by reactions with acyl and sulfonyl halides, anhydrides or isocyanates to give I (R1 = Ph; R2 = MeCO, PhCO, 4-MeC6H4SO2NHCO, 4-C1C6H4SO2NHCO).

IT 142413-14-5P 786688-43-0P 786688-43-1P
RL: SPN (Synthetic preparation); PREP (Preparation)
(preparation of (bromophenyl)aminopyrazoles)

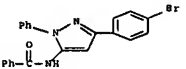
RN 342413-16-5 HCPLUS
CN Acetamide, N-[3-(4-bromophenyl)-1-phenyl-1H-pyrazol-5-yl]- (9CI) (CA INDEX NAME)



RN 786688-48-0 HCPLUS
CN Benzenesulfonamide, N-[(3-(4-bromophenyl)-1-phenyl-1H-pyrazol-5-yl)amino]carbonyl-2-chloro- (9CI) (CA INDEX NAME)



RN 786688-49-1 HCPLUS
CN Benzamide, N-[3-(4-bromophenyl)-1-phenyl-1H-pyrazol-5-yl]- (9CI) (CA INDEX NAME)



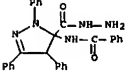
L5 ANSWER 2 OF 92 HCPLUS COPYRIGHT 2007 ACS on STN
ACCESSION NUMBER: 2003:943836 HCPLUS Full-text

DOCUMENT NUMBER: 141:243428
TITLE: Synthesis and Reactions of Some New Heterocyclic Carbohydrazides and Related Compounds as Potential Anticancer Agents
AUTHOR(S): Mansour, Abdel Kader; Eid, Mohga M.; Khalil, Nasser S. A. M.
CORPORATE SOURCE: Department of Chemistry, Faculty of Science, Cairo University, Giza, Egypt
SOURCE: Molecules (2003), 8(10), 744-755
URL: <http://www.mdpi.org/molecules/papers/81000744.pdf>

PUBLISHER: Molecular Diversity Preservation International
DOCUMENT TYPE: Journal; (online computer file)
LANGUAGE: English
OTHER SOURCE(S): CASREACT 141:243428
AB Acylation of 3-hydrazino-5,6-diphenyl-1,2,4-triazine and hydrazine hydrate with 4-aryl-1,3,7-triphenyl-8-oxa-1,2,6-triaspiro[4.4]octa-2,6-dien-9-ones gave the corresponding heterocyclic carbohydrazides. Conversion of some of the latter compds. into the versatile carbohydrazide derive. and the related oxadiazoles was undertaken. A primary in vitro test of one of the products (concentration 10-4 M) showed activity against leukemia cell lines (CCRF-CEM, K-256, MOLT-4, PRMI-8226, SR).

IT 752257-66-2P
RL: PAC (Pharmacological activity); RCT (Reactant); SPN (Synthetic preparation); BIOL (Biological study); PREP (Preparation); RACT (Reactant or reagent)
(preparation of heterocyclic carbohydrazides and related compds. as potential anticancer agents)

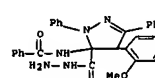
RN 752257-66-2 HCPLUS
CN 1H-Pyrazole-5-carboxylic acid, 5-(benzoylamino)-4,5-dihydro-1,3,4-triphenyl-, 2-(5,6-diphenyl-1,2,4-triazin-3-yl)hydrazide (9CI) (CA INDEX NAME)



IT 752257-67-3P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)
(preparation of heterocyclic carbohydrazides and related compds. as potential anticancer agents)

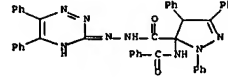
RN 752257-67-3 HCPLUS
CN 1H-Pyrazole-5-carboxylic acid, 5-(benzoylamino)-4,5-dihydro-1,3,4-triphenyl-, 2-(5,6-diphenyl-1,2,4-triazin-3-yl)hydrazide (9CI) (CA INDEX NAME)



IT 752257-64-0P 752257-65-1P 752257-68-4P
752257-65-5P 752257-70-8P

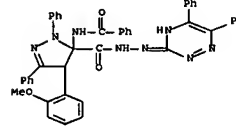
RL: SPN (Synthetic preparation); PREP (Preparation)
(preparation of heterocyclic carbohydrazides and related compds. as potential anticancer agents)

RN 752257-64-0 HCPLUS
CN 1H-Pyrazole-5-carboxylic acid, 5-(benzoylamino)-4,5-dihydro-1,3,4-triphenyl-, 2-(5,6-diphenyl-1,2,4-triazin-3-yl)hydrazide (9CI) (CA INDEX NAME)



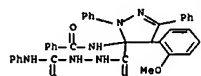
RN 752257-65-1 HCPLUS

CN 1H-Pyrazole-5-carboxylic acid, 5-(benzoylamino)-4,5-dihydro-4-(2-methoxyphenyl)-1,3-diphenyl-, 2-(5,6-diphenyl-1,2,4-triazin-3-yl)hydrazide (9CI) (CA INDEX NAME)

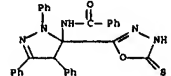


RN 752257-68-4 HCPLUS

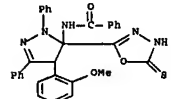
CN 1H-Pyrazole-5-carboxylic acid, 5-(benzoylamino)-4,5-dihydro-4-(2-methoxyphenyl)-1,3-diphenyl-, 2-[(phenylamino)thioxomethyl]hydrazide (9CI) (CA INDEX NAME)



RN 752257-69-5 HCAPLUS
CN Benzamide, N-[5-(4,5-dihydro-5-thioxo-1,3,4-oxadiazol-2-yl)-4,5-dihydro-1,3,4-triphenyl-1H-pyrazol-5-yl]- (9CI) (CA INDEX NAME)



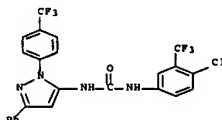
RN 752257-70-8 HCAPLUS
CN Benzamide, N-[5-(4,5-dihydro-5-thioxo-1,3,4-oxadiazol-2-yl)-4,5-dihydro-4-(2-methoxyphenyl)-1,3-diphenyl-1H-pyrazol-5-yl]- (9CI) (CA INDEX NAME)



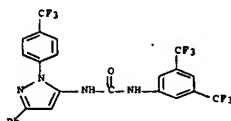
REFERENCE COUNT: 22 THERE ARE 22 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L5 ANSWER 3 OF 92 HCAPLUS COPYRIGHT 2007 ACS on STN
ACCESSION NUMBER: 2003:928892 HCAPLUS Full-text
DOCUMENT NUMBER: 140:156733
TITLE: Ureas of 5-aminopyrazole and 2-aminothiazole inhibit growth of gram-positive bacteria
AUTHOR(S): Kane, John L.; Hirth, Bradford H.; Liang, Beirong; Gourlie, Brian B.; Nahill, Sharon; Barsonian, Gary
CORPORATE SOURCE: Genzyme Drug Discovery and Development, Genzyme Corp., Cambridge, MA, 02139, USA
SOURCE: Bioorganic & Medicinal Chemistry Letters (2003), 13(24), 4463-4466
CODEN: BMCLB8; ISSN: 0960-894X
PUBLISHER: Elsevier Science B.V.
DOCUMENT TYPE: Journal
LANGUAGE: English

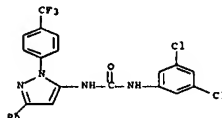
AB Ureas of 5-aminopyrazole and 2-aminothiazole emerged as lead compds. from a high-throughput screen assaying the growth of *Staphylococcus aureus*. Structure-activity relationships were developed for each compound series. Several compds. were also tested for activity against drug resistant strains of *S. aureus* in vivo.
IT 438242-75-2 ; 438242-76-3 438242-77-4
438242-92-3 656256-42-1 656256-43-0
656256-44-2 656256-45-3
RL: PAC (Pharmacological activity); BIOL (Biological study)
(ureas of 5-aminopyrazole and 2-aminothiazole inhibit growth of gram-pos. bacteria)
RN 438242-75-2 HCAPLUS
CN Urea, N-[4-chloro-3-(trifluoromethyl)phenyl]-N'-(3-phenyl-1-[4-(trifluoromethyl)phenyl]phenyl)-1H-pyrazol-5-yl]- (9CI) (CA INDEX NAME)



RN 438242-76-3 HCAPLUS
CN Urea, N-[3,5-bis(trifluoromethyl)phenyl]-N'-(3-phenyl-1-[4-(trifluoromethyl)phenyl]phenyl)-1H-pyrazol-5-yl]- (9CI) (CA INDEX NAME)

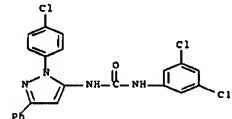


RN 438242-77-4 HCAPLUS
CN Urea, N-(3,5-dichlorophenyl)-N'-(3-phenyl-1-[4-(trifluoromethyl)phenyl]-1H-pyrazol-5-yl)- (9CI) (CA INDEX NAME)

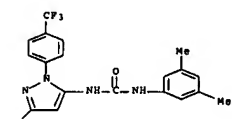


RN 438242-92-3 HCAPLUS

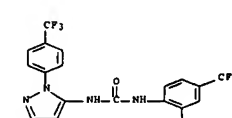
10/572,772 15/98 Robert Havlin
CN Urea, N-[1-(4-chlorophenyl)-3-phenyl-1H-pyrazol-5-yl]-N'-(3,5-dichlorophenyl)- (9CI) (CA INDEX NAME)



RN 656256-42-7 HCAPLUS
CN Urea, N-(3,5-dimethylphenyl)-N'-(3-phenyl-1-[4-(trifluoromethyl)phenyl]-1H-pyrazol-5-yl)- (9CI) (CA INDEX NAME)



RN 656256-43-8 HCAPLUS
CN Urea, N-[2-chloro-4-(trifluoromethyl)phenyl]-N'-(3-phenyl-1-[4-(trifluoromethyl)phenyl]-1H-pyrazol-5-yl)- (9CI) (CA INDEX NAME)



RN 656256-44-9 HCAPLUS
CN Urea, N-(2-methylphenyl)-N'-(3-phenyl-1-[4-(trifluoromethyl)phenyl]-1H-pyrazol-5-yl)- (9CI) (CA INDEX NAME)

AB 2-(3-Hydroxyanilino)-2-oxacetamide derivatives and interleukin 12 production inhibitors containing them
INVENTOR(S): Sato, Masakazu; Matsumura, Yuiko; Ushiki, Yasunobu; Ito, Nobumasa; Nishimura, Koji
PATENT ASSIGNEE(S): Taisho Pharmaceutical Co., Ltd., Japan
SOURCE: Jpn. Kokai Tokkyo Koho, 27 pp.
CODEN: JXKXAF
DOCUMENT TYPE: Patent
LANGUAGE: Japanese
FAMILY ACC. NUM. COUNT: 1
PATENT INFORMATION:

REFERENCE COUNT: 15 THERE ARE 15 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

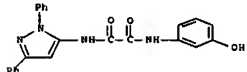
L5 ANSWER 4 OF 92 HCAPLUS COPYRIGHT 2007 ACS on STN
ACCESSION NUMBER: 2003:826823 HCAPLUS Full-text
DOCUMENT NUMBER: 139:317441
TITLE: 2-(3-Hydroxyanilino)-2-oxacetamide derivatives and interleukin 12 production inhibitors containing them

INVENTOR(S): Sato, Masakazu; Matsumura, Yuiko; Ushiki, Yasunobu; Ito, Nobumasa; Nishimura, Koji
PATENT ASSIGNEE(S): Taisho Pharmaceutical Co., Ltd., Japan
SOURCE: Jpn. Kokai Tokkyo Koho, 27 pp.
CODEN: JXKXAF

DOCUMENT TYPE: Patent
LANGUAGE: Japanese
FAMILY ACC. NUM. COUNT: 1
PATENT INFORMATION:

PATENT NO. KIND DATE APPLICATION NO. DATE
JP 2003300875 A 20031021 JP 2003-106023 20020409 --
PRIORITY APPLN. INFO.: MARPAT 139:317441
OTHER SOURCE(S): AB 3-(HOCH₂)₄NHCOCONHR [I], R = (un)substituted Ph, (un)substituted naphthyl, (un)substituted pyridyl, quinolinyl, (alkyl)benzothiophenyl, (un)substituted thiophenyl, (un)substituted pyrazolyl, substituents are given) and their pharmaceutically acceptable salts and interleukin 12 production inhibitors containing I or their salts are claimed. I [R = C₆H₃(OMe)₂-3,4] at 30 μM showed 95.7% inhibition on INF-γ-stimulated production of interleukin 12 by human peripheral blood monocytes.
IT 614722-97-3
RL: PAC (Pharmacological activity); THU (Therapeutic use); BIOL (Biological study); USES (Uses)
(preparation of 3-hydroxyanilide derivs. [N-(hetero)aryl-N'-

(hydroxyphenyl)oxalamides) as 12 production inhibitors)
 RN 614722-97-3 HCPLUS
 CN Ethenediamide, N-(1,3-diphenyl-1H-pyrazol-5-yl)-N'-(3-hydroxyphenyl)-
 (9CI) (CA INDEX NAME)



L5 ANSWER 5 OF 92 HCPLUS COPYRIGHT 2007 ACS on STN
 ACCESSION NUMBER: 20031738975 HCPLUS Full-text
 DOCUMENT NUMBER: 139:301299

TITLE: Structure-Activity Relationships of the p38α MAP Kinase Inhibitor 1-(5-tert-Butyl-2-p-tolyl-2H-pyrazol-3-yl)-3-[4-(2-morpholin-4-yl-ethoxy)naphthalen-1-yl]urea (BIRB 796)

AUTHOR(S): Regan, John; Capolino, Alison; Cirillo, Pier F.; Gilmore, Thomas; Graham, Anne G.; Hickey, Eugene; Kroes, Rachel R.; Madwed, Jeffrey; Morak, Monica; Nelson, Richard; Pargellis, Christopher A.; Swinamer, Alan; Torcellino, Carol; Tsang, Michele; Moss, Neil

CORPORATE SOURCE: Research and Development Center, Department of Medicinal Chemistry, Boehringer Ingelheim Pharmaceuticals, Ridgefield, CT, 06877, USA

SOURCE: Journal of Medicinal Chemistry (2003), 46(32), 4676-4686

PUBLISHER: American Chemical Society
 DOCUMENT TYPE: Journal
 LANGUAGE: English

OTHER SOURCE(S): CASREACT 139:301299

AB We report on the structure-activity relationships (SAR) of 1-(5-tert-butyl-2-p-tolyl-2H-pyrazol-3-yl)-3-[4-(2-morpholin-4-yl-ethoxy)naphthalen-1-yl]urea (BIRB 796), an inhibitor of p38α MAP kinase which has advanced into human clin. trials for the treatment of autoimmune diseases. Thermal denaturation was used to establish mol. binding affinities for this class of p38α inhibitors. The tert-Bu group remains a critical binding element by occupying a lipophilic domain in the kinase which is exposed upon rearrangement of the activation loop. An aromatic ring attached to N-2 of the pyrazole nucleus provides important π -CH₂ interactions with the kinase. The role of groups attached through an ethoxy group to the 4-position of the naphthalene and directed into the ATP-binding domain is elucidated. Pharmacophores with good hydrogen bonding potential, such as morpholine, pyridine, and imidazole, shift the melting temperature of p38α by 16–17° translating into Kd values of 50–100 pM. Finally, we describe several compds. that potently inhibit TNF- α production when dosed orally in mice.

IT 285983-51-9P

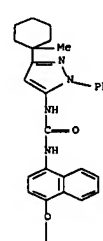
RL: PAC (Pharmacological activity); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(synthesis and p38α kinase-inhibiting activity of BIRB 796 analogs for treatment of autoimmune diseases)

RN 285983 HCPLUS

CN Urea, N-[3-(1-methylcyclohexyl)-1-phenyl-1H-pyrazol-5-yl]-N'-(4-[2-(4-

morpholinyl)ethoxy]-1-naphthalenyl]- (9CI) (CA INDEX NAME)



PAGE 1-A

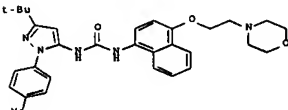
RN: GH, GM, KE, LS, MM, MZ, SD, SL, S2, T2, UC, ZM, ZW, AM, A2, BY, KG, KZ, MD, RU, TJ, TM, AT, BE, BG, CH, CY, CZ, DE, DK, EE, ES, FI, FR, GB, GR, HU, IE, IT, LU, MC, NL, PT, SE, SI, SK, TR, BP, BJ, CP, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG

AU 2003210969 A1 20030904 AU 2003-210969 20030211 --

US 2004023961 A1 20040205 US 2003-361844 20030211

PRIORITY APPLN. INFO.: US 2002-354948P P 20020211
 WO 2003-U84102 W 20030211

GI



AB 283 Of the title ureas useful for treating diseases mediated by raf kinase and diseases mediated by the VEGF induced signal transduction pathway characterized by abnormal angiogenesis or hyperpermeability processes, were claimed. Synthesis of 6 ureas such as I was described. Thus, reacting 3-(tert-butyl)-1-(4-methylphenyl)pyrazole-5-ylamine with 4-(2-morpholin-4-yl)ethoxy)naphthalen-1-yl]urea (preps. given) and CDI in CH₂Cl₂ afforded 80% I which showed IC₅₀ of < 1 μ M in in vitro raf kinase and in in vitro PIK-1 ELISA assay.

IT 285983-51-9P 285983-96-2P

RL: PAC (Pharmacological activity); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(preparation of aryl heterocycl ureas with raf kinase and angiogenesis inhibiting activity)

RN 285983-51-9 HCPLUS

CN Urea, N-[3-(1-methylcyclohexyl)-1-phenyl-1H-pyrazol-5-yl]-N'-(4-[2-(4-

morpholinyl)ethoxy]-1-naphthalenyl]- (9CI) (CA INDEX NAME)

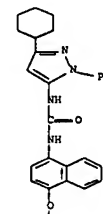


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PAGE 1-A

RN 285983-96-2 HCPLUS
 CN Urea, N-[3-(1-methylcyclohexyl)-1-phenyl-1H-pyrazol-5-yl]-N'-(4-[2-(4-

morpholinyl)ethoxy]-1-naphthalenyl]- (9CI) (CA INDEX NAME)



PAGE 1-A



PAGE 2-A

THERE ARE 17 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

ACCESSION NUMBER: 2003:633448 HCAPLUS Full-text
 DOCUMENT NUMBER: 139:185666
 TITLE: Coated pharmaceutical tablets with speckled appearance
 INVENTOR(S): Martino, Alice C.; Noack, Robert M.; Pierman, Steven A.
 PATENT ASSIGNEE(S): Pharmacia Corporation, USA
 SOURCE: PCT Int. Appl., 30 pp.
 CODEN: PIXXD2
 DOCUMENT TYPE: Patent
 LANGUAGE: English
 FAMILY ACC. NUM. COUNT: 1
 PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
WO 2003066030	A2	20030814	WO 2003-U93837	20030206 <<
WO 2003066030	A3	20031016		
W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN, CO, CR, CU, CZ, DE, DK, DM, DZ, EC, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, OM, PH, PL, PT, RO, RU, SD, SE, SG, SK, SL, TJ, TM, TN, TR, TT, TZ, UA, UG, US, UZ, VN, YU, ZA, ZM, ZW	RW: GH, GM, KE, LS, MM, MZ, SD, SL, SZ, TZ, UG, ZM, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM, AT, BE, BG, CH, CY, CZ, DE, DK, EE, ES, FI, FR, GB, GR, HU, IE, IT, LU, MC, NL, PT, SE, SI, SK, TR, BP, BJ, CF, CG, CI, CM, GA, GN, GQ, GM, ML, MR, NE, SN, TD, TG			
CA 2474921	A1	20030814	CA 2003-2474921	20030206 <<
AU 2003210930	A1	20030902	AU 2003-210930	20030206 <<
US 2003180357	A1	20030925	US 2003-359939	20030206 <<
EP 1480624	A2	20041201	EP 2003-737712	20030206
EP 1480624	B1	20041129		
R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT, IE, SI, LT, LV, PI, RO, RU, SK, TR, BG, CZ, EE, HU, SK				
BR 2003007593	A	20050201	BR 2003-7593	20030206
JP 2005517693	T	20050616	JP 2003-565454	20030206
CN 1630512	A	20050622	CN 2003-803580	20030206
NZ 533957	A	20060224	NZ 2003-533957	20030206
RU 2273473	C2	20060410	RU 2004-124065	20030206
AT 46591	T	20061215	AT 2003-737712	20030206
ZA 2004005556	A	20050810	ZA 2004-5556	20040713
NO 2004003716	A	20040906	NO 2004-3716	20040906
HK 1074581	A1	20061020	HK 2005-106918	20050811
PRIORITY APPLN. INFO.:			US 2002-356705P	P 20020207
			WO 2003-U93837	W 20030206

OTHER SOURCE(S): MARPAT 139:185666

AB A pharmaceutical tablet is provided comprising a core and a coating adherent thereto, wherein (a) the core comprises solid particles of a water-soluble dye distributed in a matrix and (b) the coating comprises gelatin gum. The tablet is suitable for peroral or intrar oral administration, for example for delivery of a drug contained in the core of the tablet to a subject. The tablet has a speckled appearance that renders the tablet readily identifiable.

IT 26170-20-1, Difenamizole
 RL: THU (Therapeutic use); BIOL (Biological study); USES (Uses)
 (active ingredients for coated pharmaceutical tablets with speckled appearance)

RN 20170-20-1 HCAPLUS
 CN Propanamide, 2-(dimethylamino)-N-(1,3-diphenyl-1H-pyrazol-5-yl)- (9CI) (CA INDEX NAME)

ACCESSION NUMBER: 2003:633447 HCAPLUS Full-text
 DOCUMENT NUMBER: 139:185665
 TITLE: Pharmaceutical dosage form for mucosal delivery
 INVENTOR(S): Martino, Alice C.; Pierman, Steven A.; Noack, Robert M.; Britten, Nancy
 PATENT ASSIGNEE(S): Pharmacia Corporation, USA
 SOURCE: PCT Int. Appl., 34 pp.
 CODEN: PIXXD2
 DOCUMENT TYPE: Patent
 LANGUAGE: English
 FAMILY ACC. NUM. COUNT: 1
 PATENT INFORMATION:

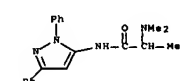
PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
WO 2003066029	A2	20030814	WO 2003-U93836	20030206 <<
WO 2003066029	A3	20031016		
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CA 2474190	A1	20030814	CA 2003-2474190	20030206 <<
AU 2003215110	A1	20030902	AU 2003-215110	20030206 <<
US 2003235617	A1	20031235	US 2003-360167	20030206 <<
EP 1471890	A2	20041103	EP 2003-710927	20030206
EP 1471890	B1	20060927		
R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT, IE, SI, LT, LV, PI, RO, RU, SK, TR, BG, CZ, EE, HU, SK				
BR 2003007473	A	20041109	BR 2003-7473	20030206
CN 1627938	A	20050615	CN 2003-803419	20030206
JP 2005519924	T	20050707	JP 2003-565453	20030206
NZ 534440	A	20060428	NZ 2003-534440	20030206
AT 340565	T	20061015	AT 2003-710927	20030206
RU 2285520	C2	20061020	RU 2004-124057	20030206
ZA 2004005614	A	20050627	ZA 2004-5614	20040714
NO 2004003723	A	20040906	NO 2004-3723	20040906
PRIORITY APPLN. INFO.:			US 2002-356705P	P 20020207
			WO 2003-U93836	W 20030206

OTHER SOURCE(S): MARPAT 139:185665

AB A pharmaceutical tablet is provided comprising an introrally disintegratable core and an excipient coating adherent thereto, wherein the coating comprises gelatin gum. The tablet is suitable for intrar oral administration, for example for delivery of a drug contained in the core of the tablet to a subject, at least in part by absorption of the drug via oral mucosa of the subject.

IT 26170-20-1, Difenamizole
 RL: THU (Therapeutic use); BIOL (Biological study); USES (Uses)

10/572,772 23/98 Robert Havlin
 (active ingredients for coated sublingual tablets)
 RN 20170-20-1 HCAPLUS
 CN Propanamide, 2-(dimethylamino)-N-(1,3-diphenyl-1H-pyrazol-5-yl)- (9CI) (CA INDEX NAME)



L5 ANSWER 9 OF 92 HCAPLUS COPYRIGHT 2007 ACS on STN
 ACCESSION NUMBER: 2003:570813 HCAPLUS Full-text
 DOCUMENT NUMBER: 139:113666
 TITLE: β -secretase inhibitors for use in treatment of diseases caused by deposits of β -amyloid peptides
 INVENTOR(S): Dietrich, Axel; Nitz, Olaf; Rester, Ulrich; Fekete, Wolfgang; Haemmerle, Marcus; Boller, Friedrich
 PATENT ASSIGNEE(S): The Genetics Company Inc., Switz.
 SOURCE: PCT Int. Appl., 42 pp.
 CODEN: PIXXD2
 DOCUMENT TYPE: Patent
 LANGUAGE: English
 FAMILY ACC. NUM. COUNT: 1
 PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
WO 2003059346	A1	20030724	WO 2003-EP504	20030120 <<
W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN, CO, CR, CU, CZ, DE, DK, DM, DZ, EC, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, OM, PH, PL, PT, RO, RU, SC, SD, SE, SG, SK, SL, TJ, TM, TN, TR, TT, TZ, UA, UG, UJ, UZ, VC, VN, YU, ZA, ZM, ZW	RW: GH, GM, KE, LS, MM, MZ, SD, SL, SZ, TZ, UG, ZM, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM, AT, BE, BG, CH, CY, CZ, DE, DK, EE, ES, FI, FR, GB, GR, HU, IE, IT, LU, MC, NL, PT, SE, SI, SK, TR, BP, BJ, CF, CG, CI, CM, GA, GN, GQ, GM, ML, MR, NE, SN, TD, TG			
CA 2473441	A1	20030724	CA 2003-2473441	20030120 <<
AU 2003205630	A1	20030730	AU 2003-205630	20030120 <<
EP 1467729	A1	20041020	EP 2003-702474	20030120
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CA 2473441	A1	20030724	CA 2003-2473441	20030120 <<
AU 2003205630	A1	20030730	AU 2003-205630	20030120 <<
EP 1467729	A1	20041020	EP 2003-702474	20030120
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JP 2005516967	T	20050609	JP 2003-559508	20030120
US 2005239999	A1	20051027	US 2005-502075	20050418
PRIORITY APPLN. INFO.:			EP 2002-1339	A 20020118
			EP 2002-12566	A 20020605
			WO 2003-EP504	W 20030120

OTHER SOURCE(S): MARPAT 139:113666

GI

10/572,772 24/98 Robert Havlin

AB The invention relates to novel substituted halophenyl inhibitors of β -secretase (II). R1 = halo, hydroxy, cyano, trifluoromethyl, Cl-4 substituted saturated or unsatd. alkyl, n-0-4; X = halo, Me, trifluoromethyl; R2 = Cl-8 alkyl containing at least one heteroatom and optionally unsatd.; R3 = aryl, carbocycle or heterocycle; R4 = R1 or a substituted aryl or heterocycle) and their use in treatment of diseases caused by deposits of β -amyloid, such as Alzheimer's disease. Thus, 7 compds. with IC50 10-170 μ M in vitro β -secretase assays are disclosed.

IT 562045-26-5
 RL: ESU (Biological study, unclassified); THU (Therapeutic use), BIOL (Biological study); USES (β -secretase inhibitors for use in treatment of diseases caused by deposits of β -amyloid peptides)
 RN 562045-26-5 HCAPLUS
 CN Cyclobutanecarboxamide, N-[3-(4-chlorophenyl)-1-(4-iodophenyl)-1H-pyrazol-5-yl]- (9CI) (CA INDEX NAME)

REFERENCE COUNT: 16 THERE ARE 16 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L5 ANSWER 10 OF 92 HCAPLUS COPYRIGHT 2007 ACS on STN
 ACCESSION NUMBER: 2003:150529 HCAPLUS Full-text
 DOCUMENT NUMBER: 138:205052
 TITLE: Preparation of 1-(pyrazol-3-yl)-3-(1-naphthyl)ureas as antiinflammatory agents
 INVENTOR(S): Cirillo, Pier Francesco; Dinallo, Roger; Regan, John Robinson; Riske, Paul S.; Swinamer, Alan David; Tan, Zhulin; Walter, Brian Andrew
 PATENT ASSIGNEE(S): Boehringer Ingelheim Pharmaceuticals, Inc., USA
 SOURCE: U.S., 44 pp., Cont.-in-part of U.S. Ser. No. 879,776, abandoned.
 CODEN: USXXAM
 DOCUMENT TYPE: Patent
 LANGUAGE: English
 FAMILY ACC. NUM. COUNT: 3
 PATENT INFORMATION:

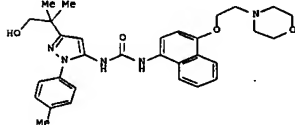
PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
US 6525046	B1	20030225	US 2002-165372	20020607 <<
US 6319921	B1	20011120	US 2000-484638	20000118 <<
US 6333325	B1	20011225	US 2001-871559	20010531 <<
US 2002058678	A1	20020516	US 2001-879776	20010612 <<
US 6329415	B1	20011211	US 2001-891579	20010626 <<
US 2002065285	A1	20020530	US 2001-891820	20010626 <<
US 6506748	B2	20030114		

PRIORITY APPLN. INFO.:

US 2000-484638 A3 20000118
US 2001-879776 B2 20010612
US 1999-116400P P 19990119

OTHER SOURCE(S): MARPAT 138:205052

GI



I

AB The title compds. Ar1NH(X)NHAr2LQ [Ar1 = pyrazolyl, pyrrolyl, imidazolyl, etc.; Ar2 = Ph, naphthyl, quinolyl, etc.; L = alkylene wherein one or more methylene groups are optionally replaced by O, N or S; Q = Ph, naphthyl, pyridyl, etc.; X = O, S], useful for treating diseases involving inflammation such as chronic inflammatory diseases, were prepared e.g., a multi-step synthesis of I, starting from Me 2,2-dimethyl-3-hydroxypropionate, was given. Representative title ureas showed IC50 of < 10 μM against TNF production in THP cells.

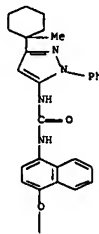
IT 285983-51-3P 285983-94-3P

RL: PAC (Pharmacological activity); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(preparation of 1-(pyrazol-3-yl)-3-(1-naphthyl)ureas as antiinflammatory agents)

RN 285983-51-9 HCAPLUS

CN Urea, N-[3-(1-methylcyclohexyl)-1-phenyl-1H-pyrazol-5-yl]-N'-(4-(2-(4-morpholinyl)ethoxy)-1-naphthalenyl)- (9CI) (CA INDEX NAME)



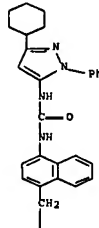
PAGE 1-A

PAGE 2-A



RN 285983-84-8 HCAPLUS
CN Urea, N-[3-(1-phenyl-1H-pyrazol-5-yl)-N'-(4-(2-(4-morpholinyl)ethoxy)-1-naphthalenyl)- (9CI) (CA INDEX NAME)

PAGE 1-A



PAGE 2-A

REFERENCE COUNT: 54 THERE ARE 54 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

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L5 ANSWER 11 OF 92 HCAPLUS COPYRIGHT 2007 ACS on STN
ACCESSION NUMBER: 2003:57886 HCAPLUS Full-text

DOCUMENT NUMBER: 138:122641

TITLE: Method of treating cytokine mediated diseases using pyrazolylureas.

INVENTOR(S): Moss, Neil; Regan, John R.

PATENT ASSIGNEE(S): Boehringer Ingelheim Pharmaceuticals, Inc., USA

SOURCE: PCT Int. Appln. 84 pp.

CODEN: PIXKD2

DOCUMENT TYPE: Patent

LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

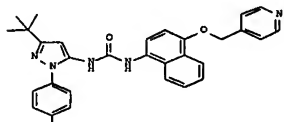
PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
WO 2003005999	A2	20030123	WO 2002-U920649	20020701 <<
WO 2003005999	A3	20030417		
WO 2003005999	A6	20040422		
W: BE, AG, AL, AM, AT, AU, AZ, BA, BB, BO, BR, BY, BZ, CA, CH, CN, CO, CR, CU, DE, DK, DM, DZ, EC, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MN, MM, MX, MZ, NO, NZ, OM, PH, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TN, TR, TT, TZ, UA, UG, UZ, VN, YU, ZA, ZM, ZW				
RW: GH, GM, KS, LB, MM, MZ, SD, SL, SZ, TZ, UG, ZM, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM, AT, BE, CH, CY, DE, DK, E9, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, TR, BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG				
CA 245316459	A1	20030123	CA 2002-2453147	20020701 <<
AU 2002316459	A1	20030129	AU 2002-316459	20020701 <<
US 2003130309	A1	20030710	US 2004-187942	20020701 <<
US 6916814	B2	20050712		
EP 1408950	A2	20040421	EP 2002-746764	20020701
EP 1408950	B1	20070125		
R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT, IE, SI, LT, LV, PI, MK, CY, AL, TR				
JP 2004536845	T	20041209	JP 2003-511806	20020701
EP 1709965	A2	20061011	EP 2006-112554	20020701
EP 1709965	A3	20061227		
R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT, IE, FI, CY, TR				
US 2004152725	A1	20040805	US 2004-761913	20040120
PRIORITY APPLN. INFO.:			US 2001-304511P	P 19990119

EP 2002-746764 A3 20020701
US 2002-187942 A3 20020701
WO 2002-U920649 W 20020701

OTHER SOURCE(S): MARPAT 138:122641

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AB A method of treating lung inflammation, endometriosis, behcet's disease, uveitis, ankylosing spondylitis, pancreatitis, cancer, percutaneous transluminal coronary angioplasty, alzheimer's disease, traumatic arthritis, sepsis, chronic obstructive pulmonary disease, and congestive heart failure comprises administration of Ar1NH(X)NHAr2LQ [Ar1 = (substituted) pyrrolyl, pyrrolidinyl, pyrazolyl, imidazolyl, oxazolyl, thiazolyl, furyl, thiényl; Ar2 = (substituted) Ph, naphthyl, quinoliny, isoquinoliny, tetrahydropyridyl, tetrahydroisoquinoliny, benzimidazolyl, benzofuryl, indanyl, indolyl, etc.; L = (o-, s-, or N-interrupted) (unsubst.) (substituted) alkylene; Q = (substituted) Ph, naphthyl, pyridyl, pyrimidinyl, imidazolyl, tetrahydropyranyl, tetrahydrafuryl dioxanyl, alkoxy, amino, etc.; X = O, S]. Thus, 5-amino-3-tart-butyl-1-(4-methylphenyl)pyrazole was stirred with COCl2 and NaHCO3 in PhMe/CH2Cl2 at 0-5° for 15 min. The organic residue was stirred overnight with 1-amino-4-(4-pyridylmethoxy)naphthalene dihydrochloride (preparation given) and diisopropylethylamine in THF to give title compound (I). Representative title compds. inhibited TNF production in THP cells with IC50<10 μM.

IT 285983-51-9P 285983-84-8P

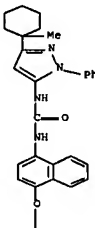
RL: PAC (Pharmacological activity); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(method of treating cytokine mediated diseases using pyrazolylureas)

RN 285983-51-9 HCAPLUS

CN Urea, N-[3-(1-methylcyclohexyl)-1-phenyl-1H-pyrazol-5-yl]-N'-(4-(2-(4-morpholinyl)ethoxy)-1-naphthalenyl)- (9CI) (CA INDEX NAME)

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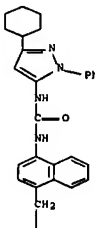
IT C285983-96-2
 RL: THU (Therapeutic use); BIOL (Biological study); USES (Uses)
 (method of treating cytokine mediated diseases using pyrazolylureas)
 RN 285983-96-2 HCAPLUS
 CN Urea, N-(3-cyclohexyl-1-phenyl-1H-pyrazol-5-yl)-N'-(4-[2-(4-morpholinyl)ethoxy]-1-naphthalenyl)- (9CI) (CA INDEX NAME)

PAGE 2-A



RN 285983-84-8 HCAPLUS
 CN Urea, N-(3-cyclohexyl-1-phenyl-1H-pyrazol-5-yl)-N'-(4-[2-(4-morpholinyl)ethyl]-1-naphthalenyl)- (9CI) (CA INDEX NAME)

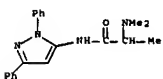
PAGE 1-A



PAGE 2-A

LS ANSWER 12 OF 92 HCAPLUS COPYRIGHT 2007 ACS on STN
 ACCESSION NUMBER: 2003:49279 HCAPLUS Full_text
 DOCUMENT NUMBER: 139:159420
 TITLE: Discrimination and selection of new potential

AUTHOR(S): antibacterial compounds using simple topological descriptors
 Murcia-Boler, Miguel; Perez-Gimenez, Facundo;
 Garcia-March, Francisco J.; Salabert-Salvador, M.
 Teresa, Diaz-Villanueva, Vladimiro; Medina-Casamayor,
 Piedad
 CORPORATE SOURCE: Faculty of Pharmacy, Department of Physical Chemistry,
 Universitat de Valencia, Valencia, Spain
 SOURCE: Journal of Molecular Graphics & Modelling (ZDOS), 21(5), 375-390
 CODEN: JMGRPF; ISSN: 1093-3263
 PUBLISHER: Elsevier Science Inc.
 DOCUMENT TYPE: Journal
 LANGUAGE: English
 AB The aim of the work was to discriminate between antibacterial and non-antibacterial drugs by topol. methods and to select new potential antibacterial agents from among new structures. The method used for antibacterial activity selection was a linear discriminant anal. (LDA). It is possible to obtain a QSAR interpretation of the information contained in the discriminant function. We make use of the pharmacol. distribution diagrams (PDDs) as a visualizing technique for the identification and selection of new antibacterial agents.
 IT 29170-20-1, Difenamicole
 RL: PAC (Pharmacological activity); THU (Therapeutic use); BIOL (Biological study); USES (Uses)
 (Discrimination and selection of new potential antibacterial compds. using simple topol. descriptors)
 RN 20170-20-1 HCAPLUS
 CN Propanamide, 2-(dimethylamino)-N-(1,3-diphenyl-1H-pyrazol-5-yl)- (9CI) (CA INDEX NAME)



REFERENCE COUNT: 20 THERE ARE 20 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

LS ANSWER 13 OF 92 HCAPLUS COPYRIGHT 2007 ACS on STN
 ACCESSION NUMBER: 2002:942809 HCAPLUS Full_text
 DOCUMENT NUMBER: 138:24709

TITLE: Preparation of pyrazole compounds and bis pyrazole-1H-pyrazole intermediates as antiinflammatory agents

INVENTOR(S): Kapadia, Suresh R.; Song, Jinhua J.; Yee, Nathan K.
 PATENT ASSIGNEE(S): Boehringer Ingelheim Pharmaceuticals, Inc., USA
 SOURCE: U.S., 37 pp., Cont.-in-part of U.S. 6,372,773.
 CODEN: USXXAM

DOCUMENT TYPE: Patent
 LANGUAGE: English
 FAMILY ACC. NUM. COUNT: 3
 PATENT INFORMATION:

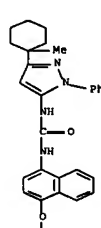
PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
US 6492529	B1	20021210	US 2002-67492	20020205 <<
US 6319921	B1	20011120	US 2000-84638	20000118 <<
US 6333325	B1	20011225	US 2001-871559	20010531 <<

10/572,772 32/98 Robert Havlin
 US 6320415 D1 20011211 US 2001-891579 20010626 <<
 US 2002065285 A1 20020530 US 2001-891820 20010626 <<
 US 6506748 B2 20030114
 US 6372773 B1 20020416 US 2001-920899 20010802 <<
 PRIORITY APPLN. INFO.: US 2000-484638 A3 20000118 <<
 US 2001-920899 A2 20010802 <<
 US 1999-116400P P 19990119 <<
 US 2001-891579 A2 20010626 <<

OTHER SOURCE(S): CASREACT 138:24709; MARPAT 138:24709
 GI

* STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY - AVAILABLE VIA OFFLINE PRINT *

AB Pyrazole compds., e.g. I, as well as bis pyrazole-1H-pyrazole intermediate compds. e.g. II, were prepared. The compds. are useful in pharmaceutical compns. for treating diseases or pathol. conditions involving inflammation such as chronic inflammatory diseases. All prepared compds. had IC50 < 10 mM for inhibition of TNF α in lipopolysaccharide stimulated THP cells.
 IT C285983-51-9 C285983-84-8P
 RL: PAC (Pharmacological activity); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)
 (preparation of pyrazole compds. and bis pyrazole-1H-pyrazole intermediates as antiinflammatory agents)
 RN 285983-51-9 HCAPLUS
 CN Urea, N-(3-(1-methylcyclohexyl)-1-phenyl-1H-pyrazol-5-yl)-N'-(4-[2-(4-morpholinyl)ethoxy]-1-naphthalenyl)- (9CI) (CA INDEX NAME)



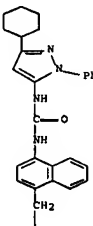
PAGE 1-A

PAGE 2-A



RN 285983-84-8 HCPLUS
CN Urea, N-(3-cyclohexyl-1-phenyl-1H-pyrazol-5-yl)-N'-(4-(2-(4-morpholinyl)ethyl)-1-naphthalenyl)- (9CI) (CA INDEX NAME)

PAGE 1-A



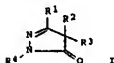
REFERENCE COUNT: 5 THERE ARE 5 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L5 ANSWER 14 OF 92 HCPLUS COPYRIGHT 2007 ACS on STN
ACCESSION NUMBER: 2002:595343 HCPLUS Full-text
DOCUMENT NUMBER: 137:150228
TITLE: Antiinflammatory compositions and methods for therapy through enhanced tissue regeneration
INVENTOR(S): Uhrich, Kathryn E.; Macedo, Braz
PATENT ASSIGNEE(S): Rutgers, The State University of New Jersey, USA
SOURCE: U.S. Pat. Appl. Publ., 17 pp.

PAGE 2-A

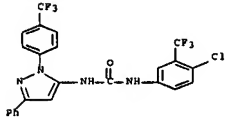


PRIORITY APPLN. INFO.: US 6410533 B1 20020625 US 2000-502101 20000210 <-
OTHER SOURCE(S): MARPAT 137:47195
GI



IT 438242-75-2P 438242-76-3P 438242-77-4P
438242-78-5P 438242-79-6P 438242-80-5P
438242-81-0P 438242-82-1P 438242-83-7P
438242-84-1P 438242-85-8P
RL: PAC (Pharmacological activity); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); US88 (Uses)
(preparation of pyrazole derivs. as antibacterial agents)

RN 438242-75-2 HCPLUS
CN Urea, N-[4-chloro-3-(trifluoromethyl)phenyl]-N'-[3-phenyl-1-[4-(trifluoromethyl)phenyl]-1H-pyrazol-5-yl]- (9CI) (CA INDEX NAME)



RN 438242-76-3 HCPLUS
CN Urea, N-[3,5-bis(trifluoromethyl)phenyl]-N'-[3-phenyl-1-[4-(trifluoromethyl)phenyl]-1H-pyrazol-5-yl]- (9CI) (CA INDEX NAME)

CODEN: USXKC0
DOCUMENT TYPE: Patent
LANGUAGE: English
FAMILY ACC. NUM. COUNT: 2
PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
US 2002106345	A1	20020808	US 2000-732516	20001207 <-
US 6648928	B2	20040203		
AU 2006201924	A1	20060601	AU 2006-201924	20060509
US 2007014832	A1	20070118	US 1999-524664	20060921
PRIORITY APPLN. INFO.:				
US 1999-304190P			P 19991207	
US 1999-455861			A 19991207	
AU 2001-19565			A 20001207	
US 2000-732516			A1 20001207	
MO 2000-US33378			A1 20001207	
US 2003-368288			BI 20030216	

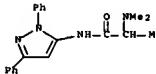
AB The invention provides methods of promoting healing through enhanced regeneration of tissue (e.g. hard tissue or soft tissue) by contacting the tissue or the surrounding tissue with an antiinflammatory agent, preferably in a controlled-release form, e.g. by dispersing the agent through a polymer matrix, appending the agent to a polymer backbone, or incorporating the agent directly into a biodegradable polymer backbone. These methods are useful in a variety of dental and orthopedic applications. Expts. are presented which demonstrate that implantation of a film comprising an aromatic polyanhydride that hydrolyzes to form a therapeutically useful salicylate resulted in less swelling in tissues adjacent to the film and a decrease in the d. of inflammatory cells as compared to other polyanhydride films. Preparation of e.g. poly[1,6-bis(o-carboxyphenoxy) hexane] is described.

IT 20170-20-1, Difemamizole

RL: PAC (Pharmacological activity); THU (Therapeutic use); BIOL (Biological study); USES (Uses)
(antiinflammatory compns. and methods for therapy through enhanced tissue regeneration)

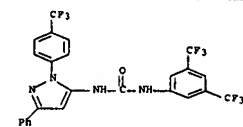
RN 20170-20-1 HCPLUS

CN Propanamide, 2-(dimethylamino)-2-(dimethylamino)-N-(1,3-diphenyl-1H-pyrazol-5-yl)- (9CI) (CA INDEX NAME)

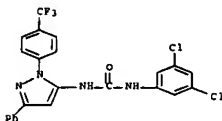


L5 ANSWER 15 OF 92 HCPLUS COPYRIGHT 2007 ACS on STN
ACCESSION NUMBER: 2002:493069 HCPLUS Full-text
DOCUMENT NUMBER: 137:47195
TITLE: Preparation of pyrazole derivs. as antibacterial agents
INVENTOR(S): Hirth, Bradford H.; Janjigian, Andrew; Vinick, Fred
PATENT ASSIGNEE(S): Genzyme Corporation, USA
SOURCE: U.S., 17 pp.
DOCUMENT TYPE: Patent
LANGUAGE: English
FAMILY ACC. NUM. COUNT: 1
PATENT INFORMATION:

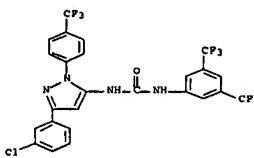
PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
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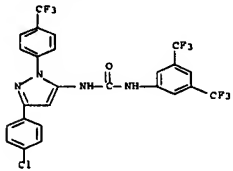
RN 438242-77-4 HCPLUS
CN Urea, N-(3,5-dichlorophenyl)-N'-[3-phenyl-1-[4-(trifluoromethyl)phenyl]-1H-pyrazol-5-yl]- (9CI) (CA INDEX NAME)



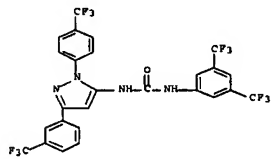
RN 438242-78-5 HCPLUS
CN Urea, N-[3,5-bis(trifluoromethyl)phenyl]-N'-[3-(3-chlorophenyl)-1-[4-(trifluoromethyl)phenyl]-1H-pyrazol-5-yl]- (9CI) (CA INDEX NAME)



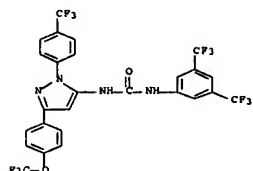
RN 438242-79-6 HCPLUS
CN Urea, N-[3,5-bis(trifluoromethyl)phenyl]-N'-[3-(4-chlorophenyl)-1-[4-(trifluoromethyl)phenyl]-1H-pyrazol-5-yl]- (9CI) (CA INDEX NAME)



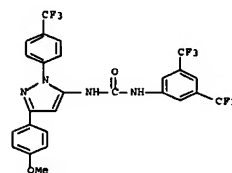
RN 438242-80-9 HCPLUS
CN Urea, N-[3,5-bis(trifluoromethyl)phenyl]-N'-(3-(trifluoromethyl)phenyl)-1-[4-(trifluoromethyl)phenyl]-1H-pyrazol-5-yl]-(9CI) (CA INDEX NAME)



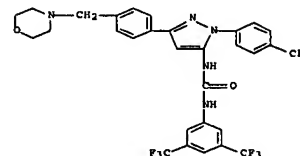
RN 438242-81-0 HCPLUS
CN Urea, N-[3,5-bis(trifluoromethyl)phenyl]-N'-(3-[4-(trifluoromethoxy)phenyl]-1-[4-(trifluoromethyl)phenyl]-1H-pyrazol-5-yl]-(9CI) (CA INDEX NAME)



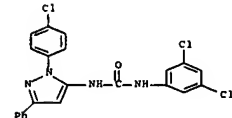
RN 438242-82-1 HCPLUS
CN Urea, N-[3,5-bis(trifluoromethyl)phenyl]-N'-(3-(4-methoxyphenyl)-1-[4-(trifluoromethyl)phenyl]-1H-pyrazol-5-yl]-(9CI) (CA INDEX NAME)



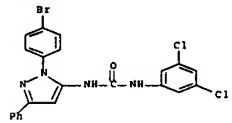
RN 438242-88-7 HCPLUS
CN Urea, N-[3,5-bis(trifluoromethyl)phenyl]-N'-(3-[4-(4-morpholinylmethyl)phenyl]-1-[4-(trifluoromethyl)phenyl]-1H-pyrazol-5-yl]-(9CI) (CA INDEX NAME)



RN 438242-92-3 HCPLUS
CN Urea, N-[1-(4-chlorophenyl)-3-phenyl-1H-pyrazol-5-yl]-N'-(3,5-dichlorophenyl)-(9CI) (CA INDEX NAME)



RN 438243-08-4 HCPLUS
CN Urea, N-[1-(4-bromophenyl)-3-phenyl-1H-pyrazol-5-yl]-N'-(3,5-dichlorophenyl)-(9CI) (CA INDEX NAME)



REFERENCE COUNT: 8 THERE ARE 8 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

LS ANSWER 16 OF 92 HCPLUS COPYRIGHT 2007 ACS on STN
ACCESSION NUMBER: 20021426876 HCPLUS Full-text
DOCUMENT NUMBER: 137:149790
TITLE: Structure-Based Classification of Antibacterial Activity
AUTHOR(S): Cronin, Mark T. D.; Aptula, Aynur O.; Dearden, John C.; Duffy, Judith C.; Netzeva, Tatiana I.; Patel, Hirani; Rowe, Philip H.; Schultz, T. Wayne; Worth, Andrew P.; Voutzoukidis, Konstantinos; Schueermann, Gerrit
CORPORATE SOURCE: School of Pharmacy and Chemistry, Liverpool John Moores University, Liverpool, L3 3AP, UK
SOURCE: Journal of Chemical Information and Computer Sciences (2002), 42(4), 869-878
CODEN: JCISD8; ISSN: 0095-2338
PUBLISHER: American Chemical Society
DOCUMENT TYPE: Journal
LANGUAGE: English
AB The aim of this study was to develop a simple quant. structure-activity relation (QSAR) for the classification and prediction of antibacterial activity, to enable *in silico* screening. To this end a database of 661 compds., classified according to whether they had antibacterial activity, and for which a total of 167 physicochem. and structural descriptors were calculated, was analyzed. To identify descriptors that allowed separation of the two classes (i.e. those compds. with and without antibacterial activity), anal. of variance was utilized and models were developed using linear discriminant and binary logistic regression analyses. Model predictivity was assessed and validated by the random removal of 30% of the compds. to form a test set, for which predictions were made from the model. The results of the analyses indicated that six descriptors, accounting for hydrophobicity and inter- and intramol. hydrogen bonding, provided excellent separation of the data. Logistic regression anal. was shown to model the data slightly more accurately than discriminant anal.
IT 23175-20-1
RL: PAC (Pharmacological activity); BIOL (Biological study)
(structure-based classification of antibacterial activity)
RN 20170-20-1 HCPLUS
CN Propanamide, 2-(dimethylamino)-N-(1,3-diphenyl-1H-pyrazol-5-yl)-(9CI) (CA INDEX NAME)

REFERENCE COUNT: 29 THERE ARE 29 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

LS ANSWER 17 OF 92 HCPLUS COPYRIGHT 2007 ACS on STN
ACCESSION NUMBER: 20021392357 HCPLUS Full-text
DOCUMENT NUMBER: 137:119059
TITLE: Pyrazole-Urea-Based Inhibitors of p38 MAP Kinase: From Lead Compound to Clinical Candidate

AUTHOR(S): Regan, John; Breitfelder, Steffani; Cirillo, Pier; Gilmore, Thomas; Graham, Anne G.; Hickey, Eugene; Klaus, Bernhard; Madewell, Jeffrey; Moriak, Monica; Moss, Neil; Pergolise, Chris; Pav, Sun; Proto, Alfred; Swinamer, Alan; Tong, Liang; Torcellini, Carol
COPARTNERS: Research and Development Center, Department of Medicinal Chemistry, Boehringer Ingelheim Pharmaceutical, Ridgefield, CT, 06877, USA

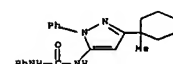
SOURCE: Journal of Medicinal Chemistry (2002), 45(14), 2994-3001
CODEN: JMCMAR; ISSN: 0022-2623
PUBLISHER: American Chemical Society

DOCUMENT TYPE: Journal
LANGUAGE: English
OTHER SOURCE(S): CASREACT 137:119059

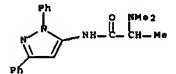
AB We report on a series of N-pyrazole, N'-aryl ureas and their mode of binding to p38 mitogen activated protein kinase. Importantly, a key binding domain that is distinct from the ATP (ATP) binding site is exposed when the conserved activation loop, consisting in part of Asp168-Gly170, adopts a conformation permitting lipophilic and hydrogen bonding interactions between this class of inhibitors and the protein. We describe the correlation of the structure-activity relationships and crystallog. structures of these inhibitors with p38. In addition, we incorporated another binding pharmacophore that forms a hydrogen bond at the ATP binding site. This modification affords significant improvements in binding, cellular, and *in vivo* potencies resulting in the selection of Compound 45 (BIRB 796) as a clin. candidate for the treatment of inflammatory diseases.

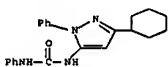
IT 223724-37-8 443913-00-6
RL: PAC (Pharmacological activity); PRP (Properties); SPN (Synthetic preparation); BIOL (Biological study)
(structure activity relationships of N-pyrazole, N'-aryl ureas and their mode of binding to p38 mitogen activated protein kinase)

RN 223724-97-8 HCPLUS
CN Urea, N-[3-(1-methylcyclohexyl)-1-phenyl-1H-pyrazol-5-yl]-N'-phenyl-(9CI) (CA INDEX NAME)



RN 443913-00-6 HCPLUS
CN Urea, N-[3-cyclohexyl-1-phenyl-1H-pyrazol-5-yl]-N'-phenyl-(9CI) (CA INDEX NAME)





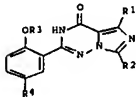
REFERENCE COUNT: 59 THERE ARE 59 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

LS ANSWER 18 OF 92 HCAPLUS COPYRIGHT 2007 ACS on STN
ACCESSION NUMBER: 2002:294376 HCAPLUS Full-text
DOCUMENT NUMBER: 137:41265
TITLE: Molecular docking and high-throughput screening for novel inhibitors of protein tyrosine phosphatase-1B
AUTHOR(S): Doman, Thompson N.; McGovern, Susan L.; Witherbee, Bryan J.; Kasten, Thomas P.; Kurumbail, Ravi; Stallings, William C.; Connolly, Daniel T.; Shochet, Brian K.
CORPORATE SOURCE: Pharmacia Corporation, Skokie, IL, 60077, USA
SOURCE: Journal of Medicinal Chemistry (2002), 45(11), 2213-2221
PUBLISHER: American Chemical Society
DOCUMENT TYPE: Journal
LANGUAGE: English
AB High-throughput screening (HTS) of compound libraries is used to discover novel leads for drug development. When a structure is available for the target, computer-based screening using mol. docking may also be considered. The two techniques have rarely been used together on the same target. The opportunity to do so presented itself in a project to discover novel inhibitors for the enzyme protein tyrosine phosphatase-1B (PTP1B), a tyrosine phosphatase that has been implicated as a key target for type II diabetes. A corporate library of approx. 400 000 compds. was screened using high-throughput exptl. techniques for compds. that inhibited PTP1B. Concurrently, mol. docking was used to screen approx. 235 000 com. available compds. against the X-ray crystallog. structure of PTP1B, and 365 high-scoring mol. were tested as inhibitors of the enzyme. Of approx. 400 000 mol. tested in the high-throughput exptl. assay, 85 (0.021%) inhibited the enzyme with IC₅₀ values less than 100 μM, the most active had an IC₅₀ value of 4.2 μM. Of the 365 mol. suggested by mol. docking, 127 (34.8%) inhibited PTP1B with IC₅₀ values less than 100 μM, the most active of these had an IC₅₀ of 1.7 μM. Structure-based docking therefore enriched the hit rate by 1700-fold over random screening. The hits from both the high-throughput and docking screens were dissimilar from phosphoryrosine, the canonical substrate group for PTP1B; the two hit lists were also very different from each other. Surprisingly, the docking hits were judged to be more druglike than the HTS hits. The diversity of both hit lists and their dissimilarity from each other suggest that docking and HTS may be complementary techniques for lead discovery.

IT 438046-45-3
RL: PAC (Pharmacological activity); PRP (Properties); BIOL (Biological study)
(mol. docking and high-throughput screening for novel inhibitors of protein tyrosine phosphatase-1B)

RN 438046-45-8 HCAPLUS

CN Urea, N-[3-(4-chlorophenyl)-1-phenyl-1H-pyrazol-5-yl]-N'-phenyl- (9CI)
(CA INDEX NAME)



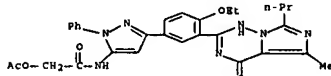
AB Title compds. [I], R1 = alkyl; R2 = cycloalkyl, alkyl; R3 = alkyl; R4 = NHSO₂RS, N(SO₂Re)SO₂R, etc., R5, R6, R7 = (substituted) vinyl, alkyl, aryl; or R5 = quinolyl, (substituted) heteraryl, etc., were prepared as phosphodiesterase I and phosphodiesterase V inhibitor (no data). Thus, 2-(5-amino-2-ethoxyphenyl)-5-methyl-7-cyclopentyl-3H-imidazo[5,1-f][1,2,4]triazin-4-one (preparation given) in THF was treated at 5° with 4-morpholinocarbonyl chloride in THF followed by stirring overnight at room temperature to give 98% 2-[2-ethoxy-5-(4-morpholinocarbonylamino)phenyl]-5-methyl-7-cyclopentyl-3H-imidazo[5,1-f][1,2,4]triazin-4-one.

IT 358390-19-99

RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); RCT (Reactant); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); RACT (Reactant or reagent); USES (Uses); (preparation of phenylimidazotriazinones as cGMP-metabolizing phosphodiesterase inhibitors)

RN 358390-19-9 HCAPLUS

CN Acetamide, 2-(acetoxyloxy)-N-[3-[3-(1,4-dihydro-5-methyl-4-oxo-7-propylimidazo[5,1-f][1,2,4]triazin-2-yl)-4-ethoxyphenyl]-1-phenyl-1H-pyrazol-5-yl]- (9CI) (CA INDEX NAME)

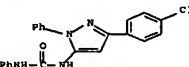


IT 358390-14-4P 358390-17-7P 358390-18-8P
358390-20-2P 358390-21-3P 358390-26-8P
259791-27-4P

RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses); (preparation of phenylimidazotriazinones as cGMP-metabolizing phosphodiesterase inhibitors)

RN 358390-14-4 HCAPLUS

CN Acetamide, N-[3-[3-(1,4-dihydro-5-methyl-4-oxo-7-propylimidazo[5,1-f][1,2,4]triazin-2-yl)-4-ethoxyphenyl]-1-phenyl-1H-pyrazol-5-yl]- (9CI) (CA INDEX NAME)



REFERENCE COUNT: 47 THERE ARE 47 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

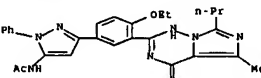
LS ANSWER 19 OF 92 HCAPLUS COPYRIGHT 2007 ACS on STN
ACCESSION NUMBER: 2001:657494 HCAPLUS Full-text
DOCUMENT NUMBER: 135:211060
TITLE: Preparation of 2-phenylimidazotriazinones as cGMP-metabolizing phosphodiesterase inhibitors
INVENTOR(S): Niewoehner, Ulrich; Es-Sayed, Mazen; Lampe, Thomas; Haning, Helmut; Schmidt, Gunther; Schlemmer, Karl-Heinz; Bischoff, Erwin; Dembowsky, Klaus; Perzborn, Elisabeth
PATENT ASSIGNEE(S): Bayer A.-G., Germany
SOURCE: Ger. Offen., 154 pp.
CODEN: GMXXBX

DOCUMENT TYPE: Patent
LANGUAGE: German
FAMILY ACC. NUM. COUNT: 1
PATENT INFORMATION:

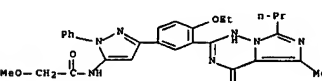
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CA 2401834		20010907	CA 2001-2401834	20010220 <-
WO 2001064677	A1	20010907	WO 2001-EP1871	20010220 <-
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EP 1280805	A1	20030205	EP 2001-911663	20010220 <-
EP 1280805	B1	20050209		
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AT 288916	T	20050215	AT 2001-911663	20010220 <-
PT 1280805	T	20050630	PT 2001-911663	20010220 <-
ES 2236190	T3	20050716	ES 2001-1911663	20010220 <-
ZA 2002006217	A	20030805	ZA 2002-6217	20020805 <-
IN 2002MH01095	A	20040529	IN 2002-MH1095	20020813
US 2004037498	A1	20040520	US 2003-220560	20030206
US 6678708	B2	20050412		
HK 1055423	A1	20060825	HK 2003-106960	20030926
US 2005267112	A1	20051201	US 2005-30605	20050105
US 7098207	B2	20060829		

PRIORITY APPLN. INFO.: DE 2000-10010067 A 20000302
WO 2001-EP1871 W 20010220
US 2003-220560 A1 20030206

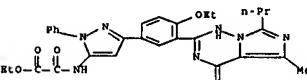
OTHER SOURCE(S): MARPAT 135:211060



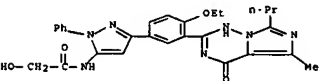
RN 358390-17-7 HCAPLUS
CN Acetamide, N-[3-[3-(1,4-dihydro-5-methyl-4-oxo-7-propylimidazo[5,1-f][1,2,4]triazin-2-yl)-4-ethoxyphenyl]-1-phenyl-1H-pyrazol-5-yl]-2-methoxy- (9CI) (CA INDEX NAME)



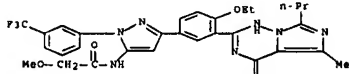
RN 358390-18-8 HCAPLUS
CN Acetic acid, [[3-[3-(1,4-dihydro-5-methyl-4-oxo-7-propylimidazo[5,1-f][1,2,4]triazin-2-yl)-4-ethoxyphenyl]-1-phenyl-1H-pyrazol-5-yl]amino]oxo- , ethyl ester (9CI) (CA INDEX NAME)



RN 358390-20-2 HCAPLUS
CN Acetamide, N-[3-[3-(1,4-dihydro-5-methyl-4-oxo-7-propylimidazo[5,1-f][1,2,4]triazin-2-yl)-4-ethoxyphenyl]-1-phenyl-1H-pyrazol-5-yl]-2-hydroxy- (9CI) (CA INDEX NAME)

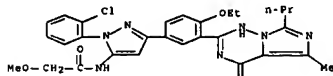


RN 358390-21-3 HCAPLUS
CN Acetamide, N-[3-[3-(1,4-dihydro-5-methyl-4-oxo-7-propylimidazo[5,1-f][1,2,4]triazin-2-yl)-4-ethoxyphenyl]-1-[3-(trifluoromethyl)phenyl]-1H-



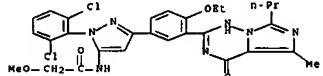
RN 358390-26-8 HCAPLUS

CN Acetamide, N-[1-(2-chlorophenyl)-3-[3-(1,4-dihydro-5-methyl-4-oxo-7-propylimidazo[5,1-f][1,2,4]triazin-2-yl)-4-ethoxyphenyl]-1H-pyrazol-5-yl]-2-methoxy- (9CI) (CA INDEX NAME)



RN 358390-27-9 HCAPLUS

CN Acetamide, N-[1-(2-dichlorophenyl)-3-[3-(1,4-dihydro-5-methyl-4-oxo-7-propylimidazo[5,1-f][1,2,4]triazin-2-yl)-4-ethoxyphenyl]-1H-pyrazol-5-yl]-2-methoxy- (9CI) (CA INDEX NAME)



LS ANSWER 20 OF 92 HCAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER: 2001:434854 HCAPLUS Full-text

DOCUMENT NUMBER: 135:51045

TITLE: Therapeutic compositions containing anti-inflammatory

agents and biodegradable polyanhydrides

INVENTOR(S): Uhrich, Kathryn; Macedo, Braz

PATENT ASSIGNEE(S): Rutgers, The State University of New Jersey, USA;

University of Medicine and Dentistry

SOURCE: PCT Int. Appl., 40 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent

LANGUAGE: English

FAMILY ACC. NUM. COUNT: 2

PATENT INFORMATION:

PATENT NO. KIND DATE APPLICATION NO. DATE

SOURCE: University of Cairo, Giza, Egypt
Indian Journal of Chemistry, Section B: Organic
Chemistry Including Medicinal Chemistry (2001
, 40B(3), 187-190

CODEN: IJSDDB; ISBN: 0376-4699

PUBLISHER: National Institute of Science Communication, CSIR

DOCUMENT TYPE: Journal

LANGUAGE: English

OTHER SOURCE(S): CASREACT 135:33441

AB The Little reaction, when carried out in chloroform in the presence of triethylamine, yields the spirocycloadducts which upon treatment with a base affords 1,3,4-triaryl-5-pyrazolecarboxamide.

IT 34346-97-3P

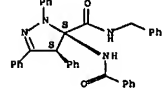
RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT
(Reactant or reagent)

(reaction of benzonitrilium N-phenylimide with (Z)-4-arylmethyleneimidazol-5(4H)-ones)

RN 34436-87-8 HCAPLUS

CN 1H-Pyrazole-5-carboxamide, 5-(benzoylamino)-4,5-dihydro-1,3,4-triphenyl-N-(phenylmethyl)-, (4R,5R)-rel- (9CI) (CA INDEX NAME)

Relative stereochemistry.



REFERENCE COUNT: 20 THERE ARE 20 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

LS ANSWER 22 OF 92 HCAPLUS COPYRIGHT 2007 ACS on STN
ACCESSION NUMBER: 2000:825371 HCAPLUS Full-text

DOCUMENT NUMBER: 134:131489

TITLE: A convenient synthesis of pyrazolo[3,4-d]pyrimidine-4,6-diones and pyrazolo[4,3-d]pyrimidine-5,7-dione derivatives

AUTHOR(S): Haddad, M. El; Soukri, M.; Lazar, S.; Benamarra, A.; Guilloumet, G.; Aksira, M.

CORPORATE SOURCE: Laboratoire de Chimie Bioorganique et Analytique, PST - Université Hassan II - Mohammedia, Mohammedia, Morocco

SOURCE: Journal of Heterocyclic Chemistry (2000),
37(5), 1247-1252

CODEN: JHTCAD; ISBN: 0022-152X

PUBLISHER: HeteroCorporation

DOCUMENT TYPE: Journal

LANGUAGE: English

OTHER SOURCE(S): CASREACT 134:131489

AB Pyrazolo[3,4-d]pyrimidine-4,6-diones and pyrazolo[4,3-d]pyrimidine-5,7-diones were synthesized by Curtius rearrangement of 3,4-pyrazoledicarboxylic acid monoesters followed by heterocyclization via urea derive. under alkaline conditions.

IT 321850-61-7P 321850-62-9P 321850-63-9P

321850-64-0P 321850-56-2P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT
(Reactant or reagent)

(preparation of pyrazolopyrimidinediones)

NO 2001041753 A2 20010614 NO 2000-US33378 20001207 <--

WO 2001041753 A3 20020912

M: AB, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN, CR, CU, CZ, DE, DK, DM, DZ, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, LZ, LR, LS, LT, LU, LV, MA, MD, MG, MK, MO, MM, MX, MZ, ND, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, TZ, UA, UG, US, UZ, VE, YU, ZA, ZW

RM: OH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, WG, ZM, AT, BR, CH, CY, DE, DK, ES, FI, PR, GB, GR, IE, IT, LU, MC, NL, PT, SE, TR, BE, BJ, CF, CG, CI, GA, GN, GW, ML, MR, NB, SN, TD, TO

CA 2393676 A1 20010614 CA 2000-2393676 20001207 <--

EP 1261347 A1 20021204 EP 2000-982544 20001207 <--

R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT, IE, SI, LT, LV, FI, RO, MK, CY, AL, TR

JP 2003528044 T 20030924 JP 2001-543098 20001207 <--

US 2004038948 A1 20040226 US 2003-366288 20030218

AU 2006201924 A1 20060601 AU 2006-201924 20060509

US 2007014632 A1 20070118 US 2006-524664 20060921

PRIORITY APPLN. INFO.: US 1999-455861 A 19991207

US 1999-304190P P 19991207

AU 2001-19565 A3 20001207

US 2000-732516 A1 20001207

WO 2000-U33378 W 20001207

US 2002-165220 B1 20020607

US 2003-368288 B1 20030218

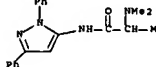
AB Methods of promoting healing through enhanced regeneration of tissue (e.g. hard tissue or soft tissue) by contacting the tissue or the surrounding tissue with an antiinflammatory agent are useful in a variety of dental and orthopedic applications. Thus, poly[1,6-bis(o-carboxyphenoxy)hexane] was prepared in a series of steps by the treatment of salicylic acid with 1,6-dibromohexane, and polymerization of the resulting 1,6-bis(o-carboxyphenoxy)hexane. The polymer was characterized by glass transition temperature measurements and then subjected to compression molding.

IT 2017-20-1, Difenamizole

RL: THU (Therapeutic use); BIOL (Biological study); USES (Uses)
(therapeutic compns. containing antiinflammatory agents and biodegradable polyanhydrides)

RN 2017-20-1 HCAPLUS

CN Propanamide, 2-(dimethylamino)-N-(1,3-diphenyl-1H-pyrazol-5-yl)- (9CI) (CA INDEX NAME)



>> d ibib abs hitstr 21-30

LS ANSWER 21 OF 92 HCAPLUS COPYRIGHT 2007 ACS on STN

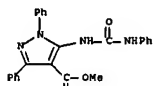
ACCESSION NUMBER: 2001:223058 HCAPLUS Full-text

DOCUMENT NUMBER: 135:33441

TITLE: Reaction of benzonitrilium N-phenylimide with (Z)-4-arylmethyleneimidazol-5(4H)-ones

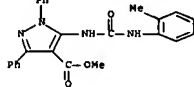
AUTHOR(S): Abdallah, M. A.; Zayed, M. E.; Shawali, A. S.

CORPORATE SOURCE: Department of Chemistry, Faculty of Science,



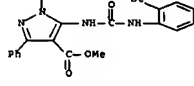
RN 321850-62-8 HCAPLUS

CN 1H-Pyrazole-4-carboxylic acid, 5-[[[(2-methylphenyl)amino]carbonyl]amino]-1,3-diphenyl-, methyl ester (9CI) (CA INDEX NAME)



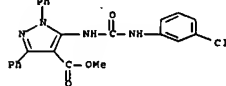
RN 321850-63-9 HCAPLUS

CN 1H-Pyrazole-4-carboxylic acid, 5-[[[(2-ethylphenyl)amino]carbonyl]amino]-1,3-diphenyl-, methyl ester (9CI) (CA INDEX NAME)

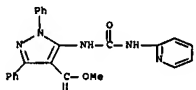


RN 321850-64-0 HCAPLUS

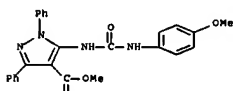
CN 1H-Pyrazole-4-carboxylic acid, 5-[[[(3-chlorophenyl)amino]carbonyl]amino]-1,3-diphenyl-, methyl ester (9CI) (CA INDEX NAME)



RN 321850-66-2 HCAPLUS
 CN 1H-Pyrazole-4-carboxylic acid, 1,3-diphenyl-5-[(2-pyridinylamino)carbonyl]amino-, methyl ester (9CI) (CA INDEX NAME)



IT 321850-65-1P
 RL: SPN (Synthetic preparation); PREP (Preparation)
 (preparation of pyrazolopyrimidinediones)
 RN 321850-65-1 HCAPLUS
 CN 1H-Pyrazole-4-carboxylic acid, 5-[[[(4-methoxyphenyl)amino]carbonyl]amino]-1,3-diphenyl-, methyl ester (9CI) (CA INDEX NAME)



REFERENCE COUNT: 28 THERE ARE 28 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L5 ANSWER 23 OF 92 HCAPLUS COPYRIGHT 2007 ACS on STN
 ACCESSION NUMBER: 2000:513688 HCAPLUS Full-text
 DOCUMENT NUMBER: 133:120325
 TITLE: Preparation of aromatic heterocyclic ureas as antinflammatory agents
 INVENTOR(S): Cirillo, Pier F.; Gilmore, Thomas A.; Hickey, Eugene R.; Regan, John R.; Zhang, Lin-Hua
 PATENT ASSIGNEE(S): Boehringer Ingelheim Pharmaceuticals, Inc., USA
 SOURCE: PCT Int. Appl., '96 pp.
 CODEN: PIKDD2

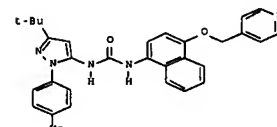
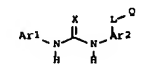
DOCUMENT TYPE: Patent
 LANGUAGE: English
 FAMILY ACC. NUM. COUNT: 3
 PATENT INFORMATION:

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WO 2000043384	A1	20000727	WO 1999-US29165	19991209 <-- W: AE, AU, BG, BR, BY, CA, CN, CZ, EE, HR, HU, ID, IL, IN, JP, KR, KZ, LT, LV, MX, NO, NZ, PL, RO, RU, SG, SI, SK, TR, UA, UZ, VN, YU, ZA

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 EP 1147104 A1 20011024 EP 1999-960668 19991209 <--
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 EE 4527 B1 20050815
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 RU 2220142 C2 20031227 RU 2001-122111 19991209 <--
 AU 770581 B2 20040236 AU 2000-17522 19991209
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 WO 1999-US29165 W 19991209
 US 2000-484638 A1 20000118

PRIORITY APPLN. INFO.:

OTHER SOURCE(S): MARPAT 133:120325
 GI



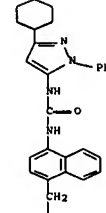
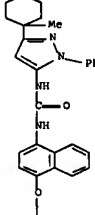
II

AB The title compds. [I; Ar1 = (un)substituted pyrrole, pyrrolidine, pyrazole, etc.; Ar2 = (un)substituted Ph, naphthyl, quinoline, etc.; L = (un)saturated (un)substituted carbon chain wherein one or more methylene groups are optionally replaced by O, N, or S; O = (un)substituted Ph, naphthyl, pyridinyl, etc.], useful in pharmaceutic compns. for

treating diseases or pathol. conditions involving inflammation such as chronic inflammatory diseases, were prepared. E.g., a multi-step synthesis of the urea II was given. Representative compds. I were evaluated and showed IC50 of < 10 μ M against TNF production in THP cells.
 IT 255983-51-9P 255983-64-8P 26592-97-2P
 RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SPN (Synthetic preparation); THU (Therapeutic use); BIO (Biological study); PREP (Preparation); USES (Uses); (preparation of aromatic heterocyclic ureas as antinflammatory agents)

RN 285983-51-9 HCAPLUS
 CN Urea, N-[3-(1-methylcyclohexyl)-1-phenyl-1H-pyrazol-5-yl]-N'-(4-[2-(4-morpholinyl)ethoxy]-1-naphthalenyl)- (9CI) (CA INDEX NAME)

PAGE 1-A



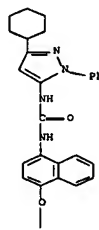
PAGE 1-A

PAGE 2-A



RN 285983-96-2 HCAPLUS
 CN Urea, N-(3-cyclohexyl-1-phenyl-1H-pyrazol-5-yl)-N'-(4-[2-(4-morpholinyl)ethyl]-1-naphthalenyl)- (9CI) (CA INDEX NAME)

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RN 285983-84-8 HCAPLUS
 CN Urea, N-(3-cyclohexyl-1-phenyl-1H-pyrazol-5-yl)-N'-(4-[2-(4-morpholinyl)ethyl]-1-naphthalenyl)- (9CI) (CA INDEX NAME)



PAGE 2-A

REFERENCE COUNT: 7 THERE ARE 7 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

LS ANSWER 24 OF 92 HCAPLUS COPYRIGHT 2007 ACS on STN
ACCESSION NUMBER: 2000457057 HCAPLUS Full-text

DOCUMENT NUMBER: 133:58801

TITLE: Preparation of aminopyrazole derivatives as p38 mitogen-activated protein (p38MAP) kinase inhibitors
Minami, Nobuyoshi; Sato, Michitaka; Hasumi, Koichi;
Yamamoto, Norio; Keino, Katayuki; Matsui, Teruaki;
Kanada, Akihiro; Ohta, Shuji; Saito, Takahisa; Sato,
Shuichiro; Asagarsu, Akira; Doi, Setsoshi; Kobayashi,
Motohiro; Sato, Jun; Asano, Hajime

PATENT ASSIGNEE(S): Teikoku Hormone Mfg. Co., Ltd., Japan
SOURCE: PCT Int. Appl., 111 pp.

CODEN: PIXKD2

DOCUMENT TYPE: Patent

LANGUAGE: Japanese

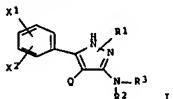
FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
WO 2000039116	A1	20000706	WO 1999-3P7186	19991221 <-
W: AU, CA, CN, JP, KR, US				
RM: AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL,				
PT, SE				
CA 2356263	A1	20000706	CA 1999-2356263	19991221 <-
EP 1142890	A1	20010110	EP 1999-559946	19991221 <-
EP 1142890	B1	20050803		
R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT,				
IE, FI				
AU 765492	B2	20030918	AU 2000-16911	19991221 <-
AT 301116	T	20050815	AT 1999-559946	19991221
ES 2244231	T3	20051201	ES 1999-559946	19991221
US 6511997	B1	20030128	US 2001-869051	20010622 <-
PRIORITY APPLN. INFO.:			JP 1998-371094	A 19981225
			WO 1999-3P7186	W 19991221

OTHER SOURCE(S): MARPAT 133:58801

GI



AB Aminopyrazole derivs. represented by general formula (I) or salts thereof [wherein X1 and X2 are each hydrogen or halogeno, or alternatively X1 and X2 may be united to form lower alkylenedioxy; Q is pyridyl or quinolyl; R1 is hydrogen or optionally substituted lower alkyl or aryl; R2 is hydrogen, lower alkyl, or aralkyl; R3 is hydrogen, an organic sulfonyl group, or -C(Y)-R4; R4 is hydrogen or an organic residue; and Y is oxygen or sulfur, with the proviso that when R3 is hydrogen, R1 is not hydrogen and R2 is hydrogen] are prepared. These compds. exhibit excellent p38MAP kinase inhibiting activities and are useful in the prevention or treatment of diseases related to tumor necrosis factor α , interleukin 1, interleukin 6 or cyclooxygenase II. These diseases include chronic articular rheumatism, multiple sclerosis, osteoarthritis, psoriasis, HIV, asthma, septic shock, inflammatory enteric disease, Crohn's disease, Alzheimer's disease, diabetes, cachexia, osteoporosis, graft vs. host disease, adult respiratory distress syndrome, arteriosclerosis, gout, glomerular nephritis (glomerulonephritis), ischemic heart failure, ulcerative colitis, septicemia, cerebral malaria, restenosis, hepatitis, systemic lupus erythematosus, thrombosis, bone resorption disease, chronic pulmonary inflammation disease, heart or kidney reperfusion disorder, cancers, Reiter's syndrome, imminent abortion, eczema, homograft rejection, seizure, fever, Behcet's disease, neuralgia, meningitis, sunburn, contact dermatitis, acute synovitis, myelitis, muscle degeneration, neovascularization, conjunctivitis, psoriatic arthritis, viral myocarditis, pancreatitis, blastoma, bleeding, arthritis, endotoxin shock, parasitic infection, tuberculosis, myocardial infarction, Hansen's disease, diabetic conjunctivitis, irritable bowel syndrome, transplant rejection, burn, bronchitis, ischemic heart disease, ectampsia, pneumonia, remission of swelling, low back pain (lumbago), myelitis, pharyngolaryngitis, Kawasaki disease, or atopic dermatitis. Thus, 300 mg Et3N was added to a suspension of 254 mg 3-amino-5-(4-fluorophenyl)-4-(4-pyridyl)pyrazone in 20 mL THF, followed by adding dropwise a solution of 464 mg phenylacetetyl chloride in 5 mL THF, and the resulting mixture was stirred at room temperature for 3 h to give 22% 5-(4-fluorophenyl)-3-phenylacetylaminol-4-(4-pyridyl)pyrazole (II). II and 5-(2-chlorophenylacetylaminol)-3-(4-fluorophenyl)-1-methyl-4-(4-pyridyl)pyrazole showed IC50 of 0.042 and 0.000088 μ g/mL against p38MAP kinase.

IT 277747-50-9P 277747-51-OP 277747-72-6P

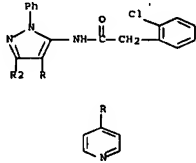
RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(preparation of aminopyrazole derivs. as p38MAP kinase inhibitors for treatment of prevention of diseases related to tumor necrosis factor α , interleukin 1, interleukin 6 or cyclooxygenase II)

RN 277747-50-9 HCAPLUS

CN Benzenacetamide, 2-chloro-N-[3-(4-fluorophenyl)-1-phenyl-4-(4-pyridinyl)-1H-pyrazol-5-yl]- α -methyl- (9CI) (CA INDEX NAME)

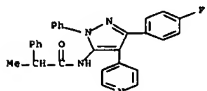
PAGE 1-A



PAGE 2-A

RN 277747-51-0 HCAPLUS

CN Benzenacetamide, N-(3-(4-fluorophenyl)-1-phenyl-4-(4-pyridinyl)-1H-pyrazol-5-yl)- α -methyl- (9CI) (CA INDEX NAME)

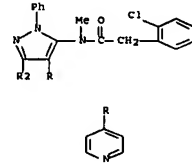


RN 277747-73-6 HCAPLUS

CN Benzenacetamide, 2-chloro-N-[3-(4-fluorophenyl)-1-phenyl-4-(4-pyridinyl)-1H-pyrazol-5-yl]-N-methyl- (9CI) (CA INDEX NAME)

PAGE 2-A

PAGE 1-A



PAGE 2-A

REFERENCE COUNT: 3 THERE ARE 3 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

LS ANSWER 25 OF 92 HCAPLUS COPYRIGHT 2007 ACS on STN
ACCESSION NUMBER: 1999699078 HCAPLUS Full-text

DOCUMENT NUMBER: 131:317778

TITLE: Phosphate derivatives for treatment of nephritis
INVENTOR(S): Miyata, Kazuyoshi; Tsuda, Yonihiko; Koji, Yasuo;
Kuroki, Morihisa; Sakai, Yasuhiro; Mukai, Kiyoishi;
Hoshimoto, Kinji; Kuri, Hideaki

PATENT ASSIGNEE(S): Otsuka Pharmaceutical Co., Ltd., Japan
SOURCE: Jpn. Kokai Tokkyo Koho, 19 pp.

CODEN: JKXXAF

DOCUMENT TYPE: Patent

LANGUAGE: Japanese

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
JP 11302177	A	19991102	JP 1998-116645	19980427 <-
PRIORITY APPLN. INFO.:			JP 1998-116645	19980427

OTHER SOURCE(S): MARPAT 131:317778

AB Phosphate derivs. (Markush structures given) are claimed for treatment of nephritis. The derivs. inhibited mesangium cell proliferation in vitro. Examples of tablets, capsules, and granules were formulated.

IT 165243-99-6

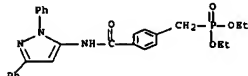
RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); THU (Therapeutic use); BIOL (Biological study); USES (Uses)

(phosphate derivs. for treatment of nephritis)

RN 169293-99-6 HCAPLUS

CN Phosphonic acid, [(4-((1,3-diphenyl-1H-pyrazol-5-

y1)amino]carbonylphenyl)methyl]-, diethyl ester (9CI) (CA INDEX NAME)

L5 ANSWER 26 OF 92 HCAPLUS COPYRIGHT 2007 ACS on STN
ACCESSION NUMBER: 1999:311199 HCAPLUS Full-text

DOCUMENT NUMBER: 130:325145

TITLE: Preparation of aromatic heterocyclic compounds as

INVENTOR(S): antinflammatory agents

Regan, John R.; Cirillo, Pier F.; Hickey, Eugene R.;
Moss, Neil; Cywin, Charles L.; Pargellis, Christopher;
Gilmore, Thomas A.

PATENT ASSIGNEE(S): Boehringen Ingelheim Pharmaceuticals, Inc., USA

SOURCE: PCT Int. Appl. '87 pp.

DOCUMENT TYPE: Patent

LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
WO 9923091	A1	19990514	WO 1998-0822907	19981029 <<
MO 9913675	A	19990524	AU 1999-13675	19981029 <<
US 6080763	A	20000627	US 1998-181743	19981029 <<
EP 1026953	A1	20000823	EP 1998-957405	19981029 <<
R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IE, IT, LU, NL, SE, MC, PT, IE, FI				
JP 2001521934	T	20011113	JP 2000-518962	19981029 <<
EP 1473292	A1	20041103	EP 2004-8840	19981029 <<
R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IE, IT, LU, NL, SE, MC, PT, IE, FI, CY				
US 6228881	B1	20010508	US 1999-461446	19991214 <<
US 2001039290	A1	20011108	US 2001-808084	20010314 <<
PRIORITY APPLN. INFO.:			US 1997-64102P	P 19971103
			EP 1998-957405	A3 19981029
			US 1998-181743	A3 19981029
			WO 1998-0822907	W 19981029
			US 1999-461446	A3 19991214

OTHER SOURCE(S): MARPAT 130:325145

GI

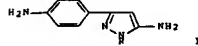
DOCUMENT NUMBER: 130:311726

TITLE: Acyl derivatives of 3-(*p*-aminophenyl)-5-aminopyrazole and its N(1)-substituted derivativesINVENTOR(S): Nam, N. L.; Grandberg, I. I.; Sorokin, V. I.
CORPORATE SOURCE: Timiryazevskoi Sel'skokhozyaistvennoi Akademii (1999), (3), 201-211SOURCE: CODEN: ITSA7; ISSN: 0021-342X
PUBLISHER: Izdatel'stvo MSKhA
DOCUMENT TYPE: Journal

LANGUAGE: Russian

OTHER SOURCE(S): CASREACT 130:311726

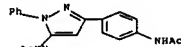
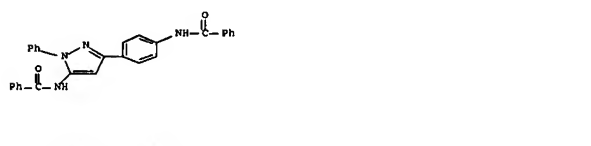
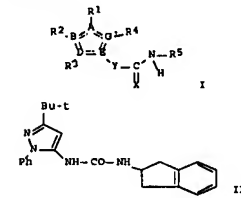
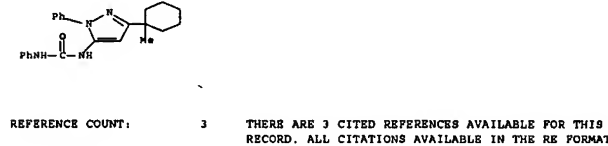
GI



AB Title compds. such as I (R = H, Me, Ph, o-tolyl, p-tolyl) were acylated on both primary amino groups.

IT 54254-75-8P 223518-55-6P 223510-59-0P
223510-61-6P 223512-65-2P
223510-74-9P 223510-76-1P 223510-90-7P
223510-85-2P 223510-87-4PRL: SPN (Synthetic preparation); PREP (Preparation)
(preparation of)

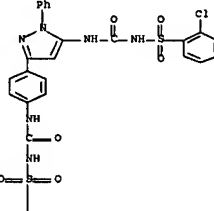
RN 94254-75-8 HCAPLUS

CN Acetamide, N-[3-(4-(acetylamino)phenyl)-1-phenyl-1H-pyrazol-5-yl]- (9CI)
(CA INDEX NAME)RN 223518-55-6 HCAPLUS
CN Benzamide, N-[3-(4-(benzoylamino)phenyl)-1-phenyl-1H-pyrazol-5-yl]- (9CI)
(CA INDEX NAME)RN 223518-59-0 HCAPLUS
CN Benzenesulfonamide, 2-chloro-N-[1-[3-[4-[[4-[(2-chlorophenyl)sulfonyl]amino]carbonyl]amino]phenyl]-1-phenyl-1H-pyrazol-5-y1]amino]carbonyl- (9CI) (CA INDEX NAME)AB The title compds. I [A = C, N; B = C, N, O, etc.; D = C, N, S; E = C, N; G = C, S, N; X = S, O, etc.; Y = NH, etc.; R1 = (un)substituted, (partially or fully halogenated) alkyl, etc., when B is C or N; R3 is Ph, naphthyl, etc., when D is C or N; or R1R2 = fused Ph or pyridinyl ring; or R2R3 = fused Ph or pyridinyl ring; R4 is H, (partially or fully halogenated) alkyl when G is C or N; R5 is Ph, naphthyl, heteroaryl, etc.] are prepared to inhibit production of cytokines involved in immunoregulation and inflammation such as interleukin-1 and tumor necrosis factor. Pyrazole derivative II was prepared from phenylhydrazine and 4,4-dimethyl-3-oxapentanenitrile. Compds. of this invention had IC50 < 10 μ M against TNF production in an *in vitro* assay using THP cells.IT 223724-97-8P
RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); BPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)
RN 223724-97-8 HCAPLUS
CN Urea, N-[3-(1-methylcyclohexyl)-1-phenyl-1H-pyrazol-5-yl]- (9CI)
(CA INDEX NAME)

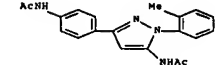
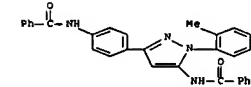
REFERENCE COUNT: 3 THERE ARE 3 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L5 ANSWER 27 OF 92 HCAPLUS COPYRIGHT 2007 ACS on STN
ACCESSION NUMBER: 1999:126025 HCAPLUS Full-text
DOCUMENT NUMBER: 130:311726
TITLE: Acyl derivatives of 3-(*p*-aminophenyl)-5-aminopyrazole and its N(1)-substituted derivativesINVENTOR(S): Nam, N. L.; Grandberg, I. I.; Sorokin, V. I.
CORPORATE SOURCE: Timiryazevskoi Sel'skokhozyaistvennoi Akademii (1999), (3), 201-211
SOURCE: CODEN: ITSA7; ISSN: 0021-342X
PUBLISHER: Izdatel'stvo MSKhA
DOCUMENT TYPE: Journal

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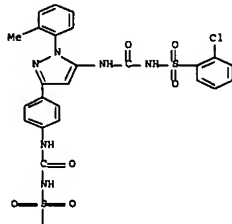


PAGE 2-A

RN 223518-63-6 HCAPLUS
CN Acetamide, N-[4-(5-(acetylamino)-1-(2-methylphenyl)-1H-pyrazol-3-yl)phenyl]- (9CI) (CA INDEX NAME)RN 223518-65-8 HCAPLUS
CN Benzamide, N-[4-(5-(benzoylamino)-1-(2-methylphenyl)-1H-pyrazol-3-yl)phenyl]- (9CI) (CA INDEX NAME)

RN 223518-69-2 HCPLUS
 CN Benzenesulfonamide, 2-chloro-N-[(4-[5-[(2-chlorophenyl)sulfonyl]amino]carbonyl)amino]-1-(2-methylphenyl)-1H-pyrazol-3-yl]phenyl]amino]carbonyl]- (9CI) (CA INDEX NAME)

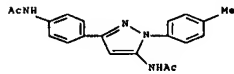
PAGE 1-A



PAGE 2-A

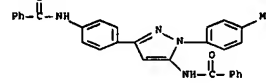


RN 223518-74-9 HCPLUS
 CN Acetamide, N-[4-(5-(acetylamino)-1-(4-methylphenyl)-1H-pyrazol-3-yl)phenyl]- (9CI) (CA INDEX NAME)



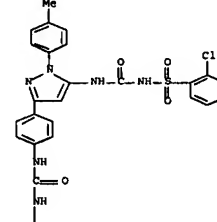
RN 223518-76-1 HCPLUS
 CN Benzamide, N-[4-(5-(benzoylamino)-1-(4-methylphenyl)-1H-pyrazol-3-yl)phenyl]- (9CI) (CA INDEX NAME)

PAGE 2-A



RN 223518-80-7 HCPLUS
 CN Benzenesulfonamide, 2-chloro-N-[(4-[5-[(2-chlorophenyl)sulfonyl]amino]carbonyl)amino]-1-(4-methylphenyl)-1H-pyrazol-3-yl]phenyl]amino]carbonyl]- (9CI) (CA INDEX NAME)

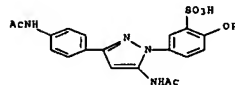
PAGE 1-A



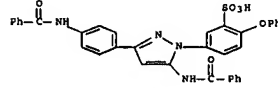
PAGE 2-A



RN 223518-85-2 HCPLUS
 CN Benzenesulfonic acid, 5-[5-(acetylamino)-3-[4-(acetylamino)phenyl]-1H-pyrazol-1-yl]-2-phenoxy- (9CI) (CA INDEX NAME)



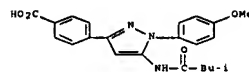
RN 223518-87-4 HCPLUS
 CN Benzenesulfonic acid, 5-[5-(benzoylamino)-3-[4-(benzoylamino)phenyl]-1H-pyrazol-1-yl]-2-phenoxy- (9CI) (CA INDEX NAME)



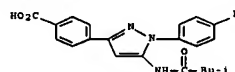
L5 ANSWER 28 OF 92 HCPLUS COPYRIGHT 2007 ACS on STN
 ACCESSION NUMBER: 1998:24217 HCPLUS Full-text
 DOCUMENT NUMBER: 128:88829
 TITLE: Solid phase synthesis of 5-aminopyrazoles and derivatives
 AUTHOR(S): Watson, Stephen P., Wilson, Richard D., Judd, Duncan B., Richards, Stephen A.
 CORPORATE SOURCE: Discovery Chemistry, Units, GlaxoWellcome Medicines Research Centre, Stevenage, SG1 2NY, UK
 SOURCE: Tetrahedron Letters (1997), 38(52), 9065-9068
 CODEN: TELRAY; ISSN: 0040-4039
 PUBLISHER: Elsevier Science Ltd.
 DOCUMENT TYPE: Journal
 LANGUAGE: English
 AB The development of a novel solid phase synthesis of some 5-aminopyrazoles and derivatives is described. Reaction of hydrazines with solid supported β -keto nitrile, 4-HO₂C₆H₄COCH₂CN, affords 5-aminopyrazoles the amino group of which is readily acylated or sulfonated. Generation of the solid supported β -keto nitrile is non-trivial and represents a key step in the overall synthesis.

IT 201139-22-2P 201139-23-3P 201139-26-6P
 201139-27-7P 201139-30-2P 201139-31-3P
 RL: SBN (Synthetic preparation); PREP (Preparation)
 (solid phase synthesis of 5-aminopyrazoles)

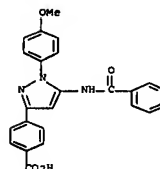
RN 201139-22-2 HCPLUS
 CN Benzoic acid, 4-[1-(4-methoxyphenyl)-5-[(3-methyl-1-oxobutyl)amino]-1H-pyrazol-3-yl]- (9CI) (CA INDEX NAME)



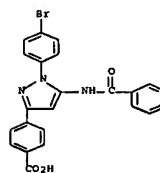
RN 201139-23-3 HCPLUS
 CN Benzoic acid, 4-[1-(4-bromophenyl)-5-[(3-methyl-1-oxobutyl)amino]-1H-pyrazol-3-yl]- (9CI) (CA INDEX NAME)



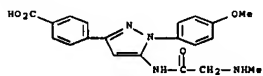
RN 201139-26-6 HCPLUS
 CN Benzoic acid, 4-[1-(4-methoxyphenyl)-5-[(3-pyridinylcarbonyl)amino]-1H-pyrazol-3-yl]- (9CI) (CA INDEX NAME)



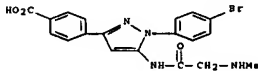
RN 201139-27-7 HCPLUS
 CN Benzoic acid, 4-[1-(4-bromophenyl)-5-[(3-pyridinylcarbonyl)amino]-1H-pyrazol-3-yl]- (9CI) (CA INDEX NAME)



RN 201139-30-2 HCPLUS
 CN Benzoic acid, 4-[1-(4-methoxyphenyl)-5-[[[(methylamino)acetyl]amino]-1H-pyrazol-3-yl]- (9CI) (CA INDEX NAME)



RN 201139-31-3 HCAPLUS
CN Benzoic acid, 4-[1-(4-bromophenyl)-5-[(methylamino)acetyl]amino]-1H-pyrazol-3-yl)- (9CI) (CA INDEX NAME)



REFERENCE COUNT: 6 THERE ARE 6 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L5 ANSWER 29 OF 92 HCAPLUS COPYRIGHT 2007 ACS on STN
ACCESSION NUMBER: 1997:579703 HCAPLUS Full-text
DOCUMENT NUMBER: 127:205576
TITLE: Preparation of sulfonylureidopyrazole derivatives as endothelin converter enzyme inhibitors
INVENTOR(S): Matsushita, Kayo; Hasegawa, Hirohiko; Kurabayashi, Yoshikazu; Ohishi, Naohito
PATENT ASSIGNEE(S): Sumitomo Pharmaceuticals Co., Ltd., Japan; Matsushita, Kayo; Hasegawa, Hirohiko; Kurabayashi, Yoshikazu; Ohishi, Naohito
SOURCE: PCT Int. Appl., 260 pp.
CODEN: PIXX2D

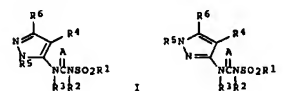
DOCUMENT TYPE: Patent
LANGUAGE: Japanese

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
WO 9730978	A1	19970828	WO 1997-JP532	19970225 <<
W: AU, CA, CN, KR, NZ, RU, US RW: AT, BE, CH, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE JP 10007658	A	19980113	JP 1997-56883	19970224 <<
CA 2247286	A1	19970828	CA 1997-2247286	19970225 <<
AU 9717354	A	19970910	AU 1997-17354	19970225 <<
EP 085890	A1	19981223	EP 1997-904634	19970225 <<
R: AT, BE, CH, DE, DK, ES, FR, GB, CR, IT, LI, NL, SE, PT, IE, PI PRIORITY APPLN. INFO.: JP 1996-65498			A 19960226	
			WO 1997-JP532	W 19970225

OTHER SOURCE(S): MARPAT 127:205576
GI



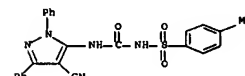
AB The title compds. (I and II; A = O, S; R1 = alkyl, alkenyl, alkynyl, cycloalkyl, etc.; R2, R3 = H, aryl, alkyl, alkenyl, alkynyl, cycloalkyl, etc.; R4, R6 = H, halo, NH2, NO2, alkyl, alkenyl, alkynyl, cycloalkyl, etc.; R5 = heterocyclyl, H, aryl, alkyl, alkenyl, alkynyl, cycloalkyl, etc.) are prepared I and II, having inhibitory effects on endothelin converter enzyme (ECE), are useful in the prevention and treatment of various circulatory disease, bronchial contraction, nervous disorder, hyposecretion, vascular lesions, various ulcers, etc. Thus, 5-amino-4-cyano-1-phenyl-(1H)-pyrazole was reacted with 4-toluenesulfonyl isocyanate to give 84.1% I (R1 = 4-MeC6H4, R2 = R3 = R6 = H, R4 = CN, R5 = Ph), which showed IC50 of 4.6 μ M against ECE.

IT 194542-43-3 HCAPLUS

RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)
(preparation of sulfonylureidopyrazole derive. as endothelin converter enzyme inhibitors)

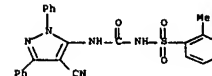
RN 194542-43-3 HCAPLUS

CN Benzenesulfonamide, N-[(4-cyano-1,3-diphenyl-1H-pyrazol-5-yl)amino]carbonyl-4-methyl- (9CI) (CA INDEX NAME)



RN 194542-44-4 HCAPLUS

CN Benzenesulfonamide, N-[(4-cyano-1,3-diphenyl-1H-pyrazol-5-yl)amino]carbonyl-2-methyl- (9CI) (CA INDEX NAME)

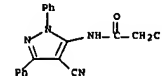


L5 ANSWER 30 OF 92 HCAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER: 1997:239113 HCAPLUS Full-text

DOCUMENT NUMBER: 126:343473

TITLE: 3-Amino-2(H)-quinolones by cyclization of N-acylated anthranilic acid derivatives

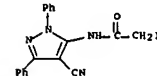


IT 189757-40-2P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)
(preparation of aminoquinolones by cyclization of N-acylated anthranilic acid derivs.)

RN 189757-40-2 HCAPLUS

CN Acetamide, N-(4-cyano-1,3-diphenyl-1H-pyrazol-5-yl)-2-iodo- (9CI) (CA INDEX NAME)



REFERENCE COUNT: 6 THERE ARE 6 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

>>
>> d ibid abs hitstr 31-50

L5 ANSWER 31 OF 92 HCAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER: 1995:860793 HCAPLUS Full-text

DOCUMENT NUMBER: 124:87730

TITLE: Synthesis of cyclodipeptides from β -pyrazolic amino acids

AUTHOR(S): El Mehdi, Ouafaa; Lavergne, Jean-Pierre; Viallefond, Philippe; Akasira, Mohamed; Sedqui, Ahmed

CORPORATE SOURCE: Lab. Amino-Acides Peptides, Univ. Montpellier II, Montpellier, 34095, Fr.

SOURCE: Bulletin de la Societe Chimique de France (1995), 132(7), 675-80

CODEN: BSCFAB; ISSN: 0037-8968

OTHER SOURCE(S): CASREACT 124:87730

AB Seven-membered ring cyclopeptides [pyrazolo[3,4-e][1,4]diazepine-4,7-diones] were prepared by a two-step procedure from β -pyrazolic amino acids.

IT 165676-73-2 172596-48-3

RL: RCT (Reactant); RACT (Reactant or reagent)
(synthesis of cyclodipeptides from pyrazolic amino acids)

RN 165676-72-2 HCAPLUS

CN 1H-Pyrazole-4-carboxylic acid, 5-[(1,1-dimethylethoxy)carbonyl]amino-1,3-diphenyl- (9CI) (CA INDEX NAME)

AUTHOR(S): Rehwald, Matthias; Gewald, Karl; Lankau, Hans-Joachim; Unverferth, Klaus

CORPORATE SOURCE: Inst. Org. Chem., Tech. Univ. Dresden, Dresden, D-01062, Germany

SOURCE: Heterocycles (1997), 45(3), 483-492

CODEN: HTCYAM; ISSN: 0385-5414

PUBLISHER: Japan Institute of Heterocyclic Chemistry

DOCUMENT TYPE: Journal

LANGUAGE: English

OTHER SOURCE(S): CASREACT 126:343473

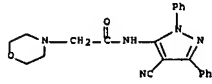
AB Reaction of secondary amines with N-(iodoacetyl)anthranilic acid derivs., 2-(iodoacetyl)acetophenone and 2-(iodoacetyl)benzophenone yielded 3-amino-2(H)-quinolones in two steps. Analogously heterocondensed 5-amino-6(H)-pyrazolo[5,4-b]pyridines were prepared. Hydroxyquinolines were subjected to Cl/OH exchange to give chloroquinolines, which are convenient for consecutive reactions.

IT 189757-41-3P 189757-42-4P

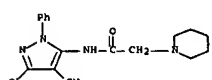
RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)
(preparation of)

RN 189757-41-3 HCAPLUS

CN 4-Morpholineacetamide, N-(4-cyano-1,3-diphenyl-1H-pyrazol-5-yl)- (9CI)
(CA INDEX NAME)



RN 189757-42-4 HCAPLUS
CN 1-Piperidineacetamide, N-(4-cyano-1,3-diphenyl-1H-pyrazol-5-yl)- (9CI)
(CA INDEX NAME)

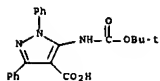


IT 189757-39-9

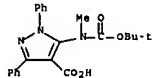
RL: RCT (Reactant); RACT (Reactant or reagent)
(preparation of aminoquinolones by cyclization of N-acylated anthranilic acid derivs.)

RN 189757-39-9 HCAPLUS

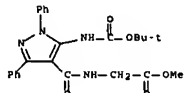
CN Acetamide, 2-chloro-N-(4-cyano-1,3-diphenyl-1H-pyrazol-5-yl)- (9CI) (CA INDEX NAME)



RN 172506-48-8 HCAPLUS
CN 1H-Pyrazole-4-carboxylic acid, 5-[(1,1-dimethylethoxy)carbonyl]methylaminoo]-1,3-diphenyl- (9CI) (CA INDEX NAME)

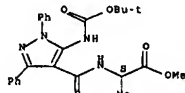


IT 172506-53-5P 172506-54-6P 172506-55-7P
172506-56-8P 172506-57-9P 172506-59-0P
RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)
(synthesis of cyclodipeptides from pyrazolic amino acids)
RN 172506-53-5 HCAPLUS
CN Glycine, N-[(5-[(1,1-dimethylethoxy)carbonyl]amino)-1,3-diphenyl-1H-pyrazol-4-yl]carbonyl-, methyl ester (9CI) (CA INDEX NAME)



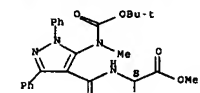
RN 172506-54-6 HCAPLUS
CN L-Alanine, N-[(5-[(1,1-dimethylethoxy)carbonyl]amino)-1,3-diphenyl-1H-pyrazol-4-yl]carbonyl-, methyl ester (9CI) (CA INDEX NAME)

Absolute stereochemistry.



RN 172506-57-9 HCAPLUS
CN L-Alanine, N-[(5-[(1,1-dimethylethoxy)carbonyl]methylamino)-1,3-diphenyl-1H-pyrazol-4-yl]carbonyl-, methyl ester (9CI) (CA INDEX NAME)

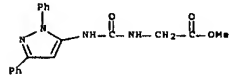
Absolute stereochemistry.



RN 172506-58-0 HCAPLUS
CN L-Phenylalanine, N-[(5-[(1,1-dimethylethoxy)carbonyl)methylamino]-1,3-diphenyl-1H-pyrazol-4-yl]carbonyl-, methyl ester (9CI) (CA INDEX NAME)

Absolute stereochemistry.

IT 172506-60-4P
RL: SPN (Synthetic preparation); PREP (Preparation)
(synthesis of cyclodipeptides from pyrazolic amino acids)
RN 172506-60-4 HCAPLUS
CN Glycine, N-[(1,3-diphenyl-1H-pyrazol-5-yl)amino]carbonyl-, methyl ester (9CI) (CA INDEX NAME)

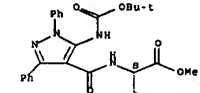


L5 ANSWER 32 OF 92 HCAPLUS COPYRIGHT 2007 ACS on STN
ACCESSION NUMBER: 1995:856442 HCAPLUS Full-text
DOCUMENT NUMBER: 123:286296
TITLE: Preparation of phosphonic diester derivatives as
antihyperlipidemics and antidiabetics
INVENTOR(S): Shoji, Yasuo; Myata, Kazuyoshi; Kuroki, Yasuhisa;
Tsuda, Yoshihiko; Tsutsumi, Kazuhiko; Inoe, Yasuhide
PATENT ASSIGNEE(S): Otsuka Pharma Co Ltd, Japan
SOURCE: Jpn. Kokai Tokkyo Koho, 13 pp.
DOCUMENT TYPE: Patent
LANGUAGE: Japanese
FAMILY ACC. NUM. COUNT: 1
PATENT INFORMATION:

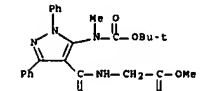
PATENT NO. KIND DATE APPLICATION NO. DATE
----- ----- ----- -----
JP 07188269 A 19950725 JP 1993-330166 19931227 --<
JP 3156026 B2 20010416 JP 1993-330166 19931227
PRIORITY APPLN. INFO.: MARPAT 123:286296
OTHER SOURCE(S): MARPAT 123:286296
AB BNHAC6H4CH2P(O)R182 [R1 = lower alkyl, Ph; A = CO, CS, SO2; B is selected from heterocyclyl (a) halo-substituted pyridine containing 1-2 of (halo-substituted) lower alkyl, CONH2, NO2, cyano, or lower alkylalkoxy]; (b) pyridine 1-oxide (containing 1-2 of (halo-substituted) lower alkyl, halo, or cyano); (c) pyrimidine containing 1-2 of lower alkyl, halo, or lower alkylthio; (d) pyrazine (containing 1-2 halo); (e) isoxazole containing 1-2 of (halo)phenyl, lower alkylphenyl, lower alkylphenyl, thiényl, phenylsulfonyl, or OH, or halo and lower alkyl; (f) pyrazole or 3-pyrazolone (containing 1-3 of lower (phenyl)alkyl, (halo)phenyl, cyano, CONH2, or thiocyanate); (g) (lower alkyl- or halo-substituted) quinoline 1-oxide; (h) 1 or 2 lower alkyl-substituted 1,8-naphthyridine] are prepared as antihyperlipidemics and antidiabetics (no data). Thus, a mixture of 3.1 g 2-amino-5-cyanopyridine-HCl and pyridine in CH2Cl2 was treated dropwise

RN 172506-55-7 HCAPLUS
CN L-Phenylalanine, N-[(5-[(1,1-dimethylethoxy)carbonyl]amino)-1,3-diphenyl-1H-pyrazol-4-yl]carbonyl-, methyl ester (9CI) (CA INDEX NAME)

Absolute stereochemistry.

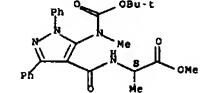


RN 172506-56-8 HCAPLUS
CN Glycine, N-[(5-[(1,1-dimethylethoxy)carbonyl]methylamino)-1,3-diphenyl-1H-pyrazol-4-yl]carbonyl-, methyl ester (9CI) (CA INDEX NAME)



RN 172506-57-9 HCAPLUS
CN L-Alanine, N-[(5-[(1,1-dimethylethoxy)carbonyl]methylamino)-1,3-diphenyl-1H-pyrazol-4-yl]carbonyl-, methyl ester (9CI) (CA INDEX NAME)

Absolute stereochemistry.



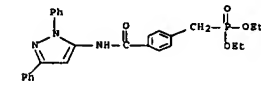
RN 172506-58-0 HCAPLUS
CN L-Phenylalanine, N-[(5-[(1,1-dimethylethoxy)carbonyl)methylamino]-1,3-diphenyl-1H-pyrazol-4-yl]carbonyl-, methyl ester (9CI) (CA INDEX NAME)

Absolute stereochemistry.

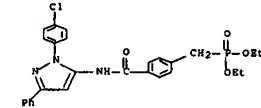
with a solution of 6.4 g 4-(diethoxyphosphoryl)methylbenzoyl chloride in CH2Cl2 under ice cooling, then treated at room temperature for 10 h to give 5.1 g diisopropyl 4-[N-(5-cyano-2-pyridyl)carbamoyl]benzylphosphonate.

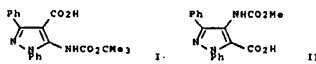
IT 169293-96-6P 169294-03-SP
RL: SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); US65 (Uses)
(preparation of heterocyclic-containing phosphonate diesters as antihyperlipidemics and antidiabetics)

RN 169293-99-6 HCAPLUS
CN Phosphonic acid, [(4-[(1,3-diphenyl-1H-pyrazol-5-yl)amino]carbonyl)phenyl]methyl-, diethyl ester (9CI) (CA INDEX NAME)



RN 169294-03-5 HCAPLUS
CN Phosphonic acid, [(4-[(4-chlorophenyl)-3-phenyl-1H-pyrazol-5-yl)amino]carbonyl)phenyl]methyl-, diethyl ester (9CI) (CA INDEX NAME)





AB Two regiosomeric pyrazolic amino acids I and II were prep'd from di-Me 1,3-diphenyl-1H-pyrazole-4,5-dicarboxylate.

IT 165676-68-0 HCPLUS

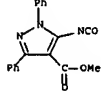
RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(preparation of regiosomeric pyrazole amino acids)

RN 165676-68-6 HCPLUS

CN 1H-Pyrazole-4-carboxylic acid, 5-isocyanato-, 1,3-diphenyl-, methyl ester

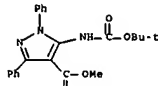
(9CI) (CA INDEX NAME)



RN 165676-70-0 HCPLUS

CN 1H-Pyrazole-4-carboxylic acid, 5-[(1,1-dimethylethoxy)carbonyl]amino)-1,3-

diphenyl-, methyl ester (9CI) (CA INDEX NAME)



IT 165676-72-2 HCPLUS

RL: SPN (Synthetic preparation); PREP (Preparation)

(preparation of regiosomeric pyrazole amino acids)

RN 165676-72-2 HCPLUS

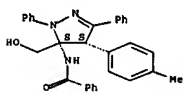
CN 1H-Pyrazole-4-carboxylic acid, 5-[(1,1-dimethylethoxy)carbonyl]amino)-1,3-

diphenyl-, methyl ester (9CI) (CA INDEX NAME)

RN 154594-05-5 HCPLUS

CN Benzamide, N-[4,5-dihydro-5-(hydroxymethyl)-4-(4-methylphenyl)-1,3-diphenyl-1H-pyrazol-5-yl]-, cis- (9CI) (CA INDEX NAME)

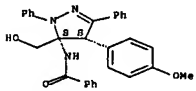
Relative stereochemistry.



RN 154594-06-6 HCPLUS

CN Benzamide, N-[4,5-dihydro-5-(hydroxymethyl)-4-(4-methoxyphenyl)-1,3-diphenyl-1H-pyrazol-5-yl]-, cis- (9CI) (CA INDEX NAME)

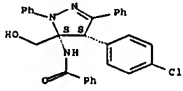
Relative stereochemistry.



RN 154594-07-7 HCPLUS

CN Benzamide, N-[4-(4-chlorophenyl)-4,5-dihydro-5-(hydroxymethyl)-1,3-diphenyl-1H-pyrazol-5-yl]-, cis- (9CI) (CA INDEX NAME)

Relative stereochemistry.



IT 129624-41-5 145383-21-7 145393-00-8

154594-02-3

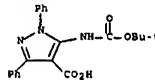
RL: RCT (Reactant); RACT (Reactant or reagent)

(reduction of)

RN 129624-41-5 HCPLUS

CN 1H-Pyrazole-5-carboxylic acid, 5-(benzoylamino)-4,5-dihydro-1,3-diphenyl-, methyl ester, cis- (9CI) (CA INDEX NAME)

Relative stereochemistry.



L5 ANSWER 34 OF 92 HCPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER: 1994:270224 HCPLUS Full-text

DOCUMENT NUMBER: 120:270224

TITLE: Synthesis and rearrangement of pyrazolylamino alcohols

AUTHOR(S): Abdallah, Magda A.; Abbas, Ikhlass M.; Mosselhi, Mosselhi A. N.; Albar, Hassan A.; Shawali, Ahmad S.

CORPORATE SOURCE: Fac. Sci., Univ. Cairo, Giza, Egypt

SOURCE: Journal of Chemical Research, Synopses (1994)

, (2), 76-7

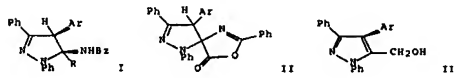
CODEN: JRPSDC; ISSN: 0308-2342

DOCUMENT TYPE: Journal

LANGUAGE: English

OTHER SOURCE(S): CASREACT 120:270224

GI



AB 4-Aryl-5-benzoylamino-5-hydroxymethyl-1,3-diphenyl-2-pyrazolines I (R = CH₂OH, Ar = Ph, 4-MeC₆H₄, 4-MeOC₆H₄, 4-ClC₆H₄) were prepared by LiAlH₄ reduction of either the spiropyrazolines II or the corresponding pyrazoline esters I (R = CO₂Me); treatment of I (R = CH₂OH) with hydrochloric acid in dioxane at room temperature, gave 4-aryl-5-hydroxymethyl-1,3-diphenylpyrazoles III.

IT 154594-04-4P 154594-05-5P 154594-06-6P

1E4E54-07-7P

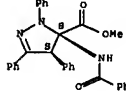
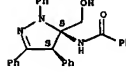
RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(preparation and rearrangement of)

RN 154594-04-4 HCPLUS

CN Benzamide, N-[4,5-dihydro-5-(hydroxymethyl)-1,3,4-triphenyl-1H-pyrazol-5-yl]-, cis- (9CI) (CA INDEX NAME)

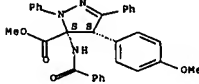
Relative stereochemistry.



RN 145383-21-7 HCPLUS

CN 1H-Pyrazole-5-carboxylic acid, 5-(benzoylamino)-4,5-dihydro-4-(4-methoxyphenyl)-1,3-diphenyl-, methyl ester, cis- (9CI) (CA INDEX NAME)

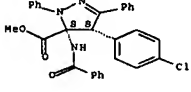
Relative stereochemistry.



RN 145383-22-8 HCPLUS

CN 1H-Pyrazole-5-carboxylic acid, 5-(benzoylamino)-4-(4-chlorophenyl)-4,5-dihydro-1,3-diphenyl-, methyl ester, cis- (9CI) (CA INDEX NAME)

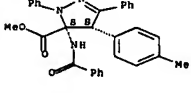
Relative stereochemistry.



RN 154594-03-3 HCPLUS

CN 1H-Pyrazole-5-carboxylic acid, 5-(benzoylamino)-4-(4-methylphenyl)-1,3-diphenyl-, methyl ester, cis- (9CI) (CA INDEX NAME)

Relative stereochemistry.

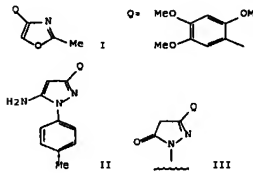


10/572,772

77/98

Robert Havlin

ACCESSION NUMBER: 1994:217403 HCAPLUS Full-text
 DOCUMENT NUMBER: 120:217403
 TITLE: Synthesis and spectroscopy of new substituted arylazoles
 AUTHOR(S): Sanchez-Viecas, F.; Gomez, Maria R.
 CORPORATE SOURCE: Fac. Quim., UNAM, Mexico City, 04510, Mex.
 SOURCE: Revista Latinoamericana de Quimica (1991), 22(3), 85-9
 CODEN: RLQAQ8; ISSN: 0370-5943
 DOCUMENT TYPE: Journal
 LANGUAGE: Spanish
 OTHER SOURCE(S): CASREACT 120:217403
 GI



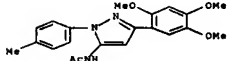
AB Methyl(trimethoxyphenyl)oxazole I could not be prepared by cyclocondensation of an α -halo ketone and an amide, but was prepared in 72% yield from acyloin ester $\text{OCOCH}_2\text{COAc}$ and NH_2OAc in AcOH. It could not be converted to its imidazole analog, which also could not be prepared by other routes. This is presumably due to special reactivity of such 2,4,5-trimethoxyphenyl compds., as seen previously. On the other hand, cyclization of OCOCH_2CN with p-MeC₆H₄NH₂·HCl gave 70% aminopyrazole derivative II, which was N-acetylated in 74% yield. Similar cyclization of $\text{OCOCH}_2\text{CONH}_2$ gave 97% pyrazolinone III. Some characteristic IR, 1H-NMR, and mass spectral data are given and discussed.

IT 153981-93-2P

RL: SPN (Synthetic preparation); PREP (Preparation)
(preparation of arylazoles and arylpyrazoles)

RN 153981-93-2 HCAPLUS

CN Acetamide, N-[1-(4-methoxyphenyl)-3-(2,4,5-trimethoxyphenyl)-1H-pyrazol-5-yl]- (9CI) (CA INDEX NAME)

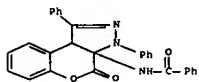
LS ANSWER 36 OF 92 HCAPLUS COPYRIGHT 2007 ACS on STN
ACCESSION NUMBER: 1994:30710 HCAPLUS Full-textDOCUMENT NUMBER: 120:30710
TITLE: Synthesis of aminopyrazolecarboxylic acid derivatives

10/572,772

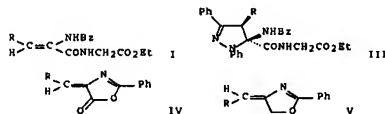
79/98

Robert Havlin

RN 151806-43-8 HCAPLUS
 CN Benzamide, N-(3,5b-dihydro-4-oxo-1,3-diphenyl[1]benzopyrano[3,4-c]pyrazol-3a(4H)-yl)- (9CI) (CA INDEX NAME)



LS ANSWER 37 OF 92 HCAPLUS COPYRIGHT 2007 ACS on STN
ACCESSION NUMBER: 1993:581191 HCAPLUS Full-text
 DOCUMENT NUMBER: 119:181191
 TITLE: 1,3-Dipolar cycloaddition of benzonitrilium N-phenylimide to didehydropeptides
 AUTHOR(S): Abdallah, Magda A.; Albar, Hassan A.; Shawali, Ahmad S.
 CORPORATE SOURCE: Fac. Sci., Univ. Cairo, Giza, Egypt
 JOURNAL: Journal of Chemical Research, Synopses (1993), (S), 182-3
 CODEN: JRPSDC; ISSN: 0308-2342
 DOCUMENT TYPE: Journal
 LANGUAGE: English
 OTHER SOURCE(S): CASREACT 119:181191
 GI



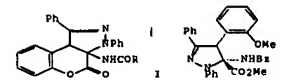
AB Didehydropeptides I ($\text{Ar} = \text{Ph}, 4\text{-MeC}_6\text{H}_4, 4\text{-ClC}_6\text{H}_4, 4\text{-O}_2\text{NC}_6\text{H}_4, 3,4\text{-methylene dioxyphenyl}$) underwent a regioselective 1,3-dipolar cycloaddn. reaction with benzonitrilium N-phenylimide PhC₆H₄N+Ph (II) to give cycloadducts III. I were obtained by the ring cleavage of (Z)-oxazolones IV or (E)-oxazolones V with H-Gly-OEt·HCl in the presence of Et₃N in DMP. II was generated in situ from PhC(=NNHPh)Cl by treatment with Et₃N.

10/572,772

78/98

Robert Havlin

AUTHOR(S): Shawali, Ahmad S.; Hassaneen, Hamdi M.; Albert, Hassan A.; Abdelhamid, Hyam A.
 CORPORATE SOURCE: Fac. Sci., Univ. Cairo, Giza, Egypt
 SOURCE: Indian Journal of Chemistry, Section B: Organic Chemistry Including Medicinal Chemistry (1992), 32B(7), 795-6
 DOCUMENT TYPE: Journal
 LANGUAGE: English
 OTHER SOURCE(S): CASREACT 120:30710
 GI



AB Diphenylnitrilimine adds regioselectively to 3-acylaminocoumarins and Me α -acylamino-o-methoxycinnamate to yield exclusively the cycloadducts I (R = Me, Ph) and II, resp. I were converted into II by their treatment with KOH and di-Me sulfate in methanol. The regiochem. of the cycloadducts have been confirmed by their conversion to the known 1,3-diphenylchromeno[3,4-c]pyrazol-4-(3H)-one and Me 1,3-diphenyl-4-(o-methoxyphenyl)-pyrazole-5-carboxylate, resp.

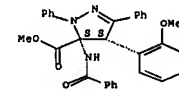
IT 14533-20-6P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

RN 14533-20-6 HCAPLUS

CN 1H-Pyrazole-5-carboxylic acid, 5-(benzoylamo)-4,5-dihydro-4-(2-methoxyphenyl)-1,3-diphenyl-, methyl ester, cis- (9CI) (CA INDEX NAME)

Relative stereochemistry.



IT 151806-42-7P 151806-43-8P

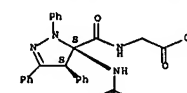
RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

RN 151806-42-7 HCAPLUS

CN Acetamide, N-(3,5b-dihydro-4-oxo-1,3-diphenyl[1]benzopyrano[3,4-c]pyrazol-3a(4H)-yl)- (9CI) (CA INDEX NAME)

IT 150330-79-3P 150330-80-6P 150330-81-7P
 150330-82-8P 150330-83-9P 150330-64-OPRL: SPN (Synthetic preparation); PREP (Preparation)
(preparation and ring cleavage of)
 RN 150330-79-3 HCAPLUS
 CN Glycine, N-[5-(benzoylamo)-4,5-dihydro-4-oxo-1,3,4-triphenyl-1H-pyrazol-5-yl]carbonyl-, ethyl ester, cis- (9CI) (CA INDEX NAME)

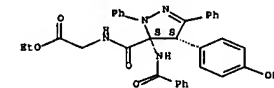
Relative stereochemistry.



RN 150330-80-6 HCAPLUS

CN Glycine, N-[5-(benzoylamo)-4,5-dihydro-4-(4-methoxyphenyl)-1,3-diphenyl-1H-pyrazol-5-yl]carbonyl-, ethyl ester, cis- (9CI) (CA INDEX NAME)

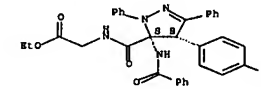
Relative stereochemistry.



RN 150330-81-7 HCAPLUS

CN Glycine, N-[5-(benzoylamo)-4,5-dihydro-4-(4-methylphenyl)-1,3-diphenyl-1H-pyrazol-5-yl]carbonyl-, ethyl ester, cis- (9CI) (CA INDEX NAME)

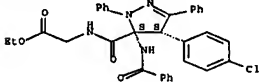
Relative stereochemistry.



RN 150330-82-8 HCAPLUS

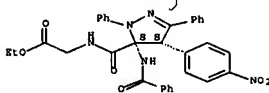
CN Glycine, N-[5-(benzoylamo)-4-(4-chlorophenyl)-4,5-dihydro-1,3-diphenyl-1H-pyrazol-5-yl]carbonyl-, ethyl ester, cis- (9CI) (CA INDEX NAME)

Relative stereochemistry.



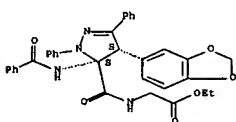
RN 150330-83-9 HCAPLUS
CN Glycine, N-[5-(benzoylamino)-4,5-dihydro-4-(4-nitrophenyl)-1,3-diphenyl-1H-pyrazol-5-yl]carbonyl-, ethyl ester, cis- (9CI) (CA INDEX NAME)

Relative stereochemistry.

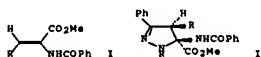


RN 150330-84-0 HCAPLUS
CN Glycine, N-[4-(1,3-benzodioxol-5-yl)-5-(benzoylamino)-4,5-dihydro-1,3-diphenyl-1H-pyrazol-5-yl]carbonyl-, ethyl ester, cis- (9CI) (CA INDEX NAME)

Relative stereochemistry.



L5 ANSWER 38 OF 92 HCAPLUS COPYRIGHT 2007 ACS on STN
ACCESSION NUMBER: 1992:59626 HCAPLUS Full-text
DOCUMENT NUMBER: 118:59626
TITLE: 1,3-Dipolar cycloaddition reactions of diphenylnitrilimine with esters of α,β -didehydro amino acids
AUTHOR(S): Shawali, Ahmad S.; Fahmi, Abdellgawad A.; Hassaneen, Hamsi M.; Abdallah, Magda A.; Abdelhamid, Hyam A.
CORPORATE SOURCE: Fac. Sci., Univ. Cairo, Giza, Egypt
SOURCE: Journal of Chemical Research, Synopses (1992), (11), 360-1
CODEN: JRPBDC; ISSN: 0308-2342
DOCUMENT TYPE: Journal
LANGUAGE: English
OTHER SOURCE(S): CASREACT 118:59626
GI

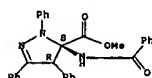


AB Diphenylnitrilimine adds regioselectively to the carbon-carbon double bond of Me β -aryl-N-benzoyl- α,β -didehydroalaninates I (R = aryl) to afford the pyrazoline derivs. II (R = aryl). The regiochem. of the latter cycloadducts was evidenced chemical by their conversion to the known 1,3,4-triarylpolyazoles-5-carboxylates and by their alternative synthesis from the recently reported spiropyrazoles. The 1H NMR spectral data of II were compatible with their assigned structure.

IT 139285-17-9F 145382-27-3P
RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent); (preparation and attempted thermolysis and isomerization of)

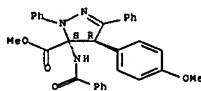
RN 139285-17-9 HCAPLUS
CN 1H-Pyrazole-5-carboxylic acid, 5-(benzoylamino)-4,5-dihydro-1,3,4-triphenyl-, methyl ester, trans- (9CI) (CA INDEX NAME)

Relative stereochemistry.



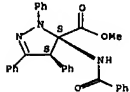
RN 145383-27-3 HCAPLUS
CN 1H-Pyrazole-5-carboxylic acid, 5-(benzoylamino)-4,5-dihydro-4-(4-methoxyphenyl)-1,3-diphenyl-, methyl ester, trans- (9CI) (CA INDEX NAME)

Relative stereochemistry.



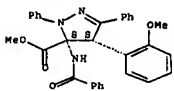
IT 139624-41-5P 145383-20-6P
RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent); (preparation and methanolysis of)
RN 129624-41-5 HCAPLUS
CN 1H-Pyrazole-5-carboxylic acid, 5-(benzoylamino)-4,5-dihydro-1,3,4-triphenyl-, methyl ester, cis- (9CI) (CA INDEX NAME)

Relative stereochemistry.



RN 145383-20-6 HCAPLUS
CN 1H-Pyrazole-5-carboxylic acid, 5-(benzoylamino)-4,5-dihydro-4-(2-methoxyphenyl)-1,3-diphenyl-, methyl ester, cis- (9CI) (CA INDEX NAME)

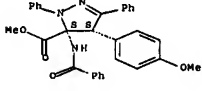
Relative stereochemistry.



IT 145383-21-7P 145383-21-6P 145383-21-5P
145383-24-CP
RL: SPN (Synthetic preparation); PREP (Preparation); (preparation of, by regioselective and stereoselective cycloaddn. of diphenylnitrilimine with Me β -aryl-N-benzoyl- α,β -didehydroalaninate)

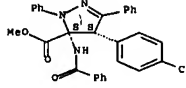
RN 145383-21-7 HCAPLUS
CN 1H-Pyrazole-5-carboxylic acid, 5-(benzoylamino)-4,5-dihydro-4-(4-methoxyphenyl)-1,3-diphenyl-, methyl ester, cis- (9CI) (CA INDEX NAME)

Relative stereochemistry.



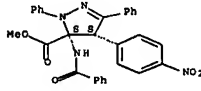
RN 145383-22-8 HCAPLUS
CN 1H-Pyrazole-5-carboxylic acid, 5-(benzoylamino)-4-(4-chlorophenyl)-4,5-dihydro-1,3-diphenyl-, methyl ester, cis- (9CI) (CA INDEX NAME)

Relative stereochemistry.



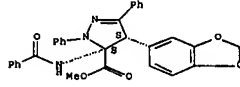
RN 145383-23-9 HCAPLUS
CN 1H-Pyrazole-5-carboxylic acid, 5-(benzoylamino)-4,5-dihydro-4-(4-nitrophenyl)-1,3-diphenyl-, methyl ester, cis- (9CI) (CA INDEX NAME)

Relative stereochemistry.



RN 145383-24-0 HCAPLUS
CN 1H-Pyrazole-5-carboxylic acid, 4-(1,3-benzodioxol-5-yl)-5-(benzoylamino)-4,5-dihydro-1,3-diphenyl-, methyl ester, cis- (9CI) (CA INDEX NAME)

Relative stereochemistry.



L5 ANSWER 39 OF 92 HCAPLUS COPYRIGHT 2007 ACS on STN
ACCESSION NUMBER: 1992:128760 HCAPLUS Full-text
DOCUMENT NUMBER: 116:128760
TITLE: On the synthesis of geminally functionalized heterocyclic aminocarboxylic acid esters
AUTHOR(S): Coutoulis-Arygiropoulos, E.; Thessalonikeos, E.
CORPORATE SOURCE: Lab. Org. Chem., Univ. Thessaloniki, Thessaloniki, 540 06, Greece
SOURCE: Journal of Heterocyclic Chemistry (1992), 28 (8), 1945-8
CODEN: JHTCAD; ISSN: 0022-152X
DOCUMENT TYPE: Journal
LANGUAGE: English
OTHER SOURCE(S): CASREACT 116:128760
GI



AB The title compds. I ($X = O$, Ar = mesityl, 2,6-C₁₂C₆H₃; X = NPh, Ar = Ph, 4-MeC₆H₄, 4-C₁₂C₆H₄) were easily prepared by two alternative procedures: 1,3-dipolar cycloaddn. to benzamidocinamates prepared by methanolysis of the corresponding oxazolones or methanolysis of the spirooxazolones II, synthesized by 1,3-dipolar cycloaddn. to oxazolones. Both reaction sequences show the same stereo- and regioselectivity.

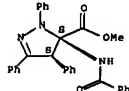
IT 139224-41-5
RL: RCT (Reactant); **SPN** (Synthetic preparation); **PREP** (Preparation); **RACT** (Reactant or reagent)

(preparation and aromatization of)

RN 129624-41-5 HCPLUS

CN 1H-Pyrazole-5-carboxylic acid, 5-(benzoylamino)-4,5-dihydro-1,3,4-triphenyl-, methyl ester, cis- (9CI) (CA INDEX NAME)

Relative stereochemistry.

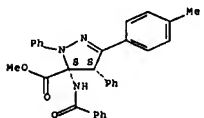


IT 139225-15-7P 139225-16-aP 139225-17-9P
RL: SPN (Synthetic preparation); PREP (Preparation)
 (preparation of)

RN 139225-15-7 HCPLUS

CN 1H-Pyrazole-5-carboxylic acid, 5-(benzoylamino)-4,5-dihydro-3-(4-methylphenyl)-1,4-diphenyl-, methyl ester, cis- (9CI) (CA INDEX NAME)

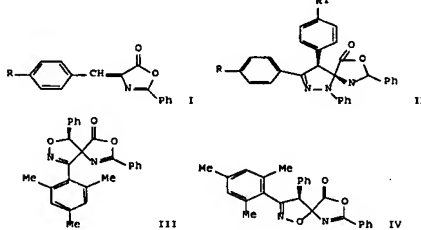
Relative stereochemistry.



RN 139225-16-8 HCPLUS

CN 1H-Pyrazole-5-carboxylic acid, 5-(benzoylamino)-3-(4-chlorophenyl)-4,5-dihydro-1,4-diphenyl-, methyl ester, cis- (9CI) (CA INDEX NAME)

Relative stereochemistry.



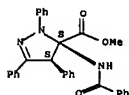
AB The title reaction of phenylarylideneoxazolones I ($R = H, Me, MeO$) with 4-RIC₆H₄C₁Ph-N,N-Ph (R1 = H, Me, Cl), generated from 4-RIC₆H₄COCl:NHPh with Et₃N, gave spiro[pyrazoline-oxazolinones] II. The structure of II was supported by anal. and spectral data. The regiochem. of these cycloaddns. suggested that nitrile oxides might add in a similar fashion, thus, the nitrile oxide-cycloadduct previously assigned structure III was reexamined and shown to be the regioisomer IV.

IT 139224-41-2P 129624-42-6P
RL: SPN (Synthetic preparation); PREP (Preparation)
 (preparation of)

RN 129624-41-5 HCPLUS

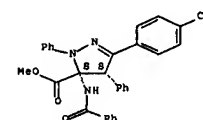
CN 1H-Pyrazole-5-carboxylic acid, 5-(benzoylamino)-4,5-dihydro-1,3,4-triphenyl-, methyl ester, cis- (9CI) (CA INDEX NAME)

Relative stereochemistry.



RN 129624-42-6 HCPLUS
CN 1H-Pyrazole-5-carboxamide, 5-(benzoylamino)-4,5-dihydro-N-(4-methylphenyl)-1,3,4-triphenyl-, cis- (9CI) (CA INDEX NAME)

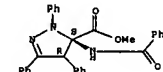
Relative stereochemistry.



RN 139205-17-9 HCPLUS

CN 1H-Pyrazole-5-carboxylic acid, 5-(benzoylamino)-4,5-dihydro-1,3,4-triphenyl-, methyl ester, trans- (9CI) (CA INDEX NAME)

Relative stereochemistry.



**L5 ANSWER 40 OF 92 HCPLUS COPYRIGHT 2007 ACS on STN
 ACCESSION NUMBER: 1990:552311 HCPLUS Full-text**

DOCUMENT NUMBER: 113:152311

TITLE: 1,3-Dipolar Cycloaddition reactions of 2-phenyl-4-arylideneoxazol-5(NH)-ones with nitrile imines. A reinvestigation of the regiochemistry of the 1,3-dipolar cycloaddition reactions of 2-phenyl-4-arylideneoxazol-5(NH)-ones with nitrile oxides

AUTHOR(S): Coutouli-Argepoulou, Eudoxia; Argyropoulos, Nikolaos G.; Thessalonikeos, Elias

CORPORATE SOURCE: Dep. Chem., Univ. Thessaloniki, Thessaloniki, 54006, Greece

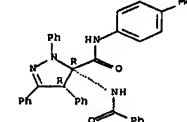
SOURCE: Journal of Chemical Research, Synopses (1990), (7), 202-3

DOCUMENT TYPE: CODEN: JRPSCD; ISSN: 0308-2342

LANGUAGE: English

OTHER SOURCE(S): CASREACT 113:152311

GI



**L5 ANSWER 41 OF 92 HCPLUS COPYRIGHT 2007 ACS on STN
 ACCESSION NUMBER: 1989:534747 HCPLUS Full-text**

DOCUMENT NUMBER: 111:134747

TITLE: Preparation and testing of heterocycliccarboxylglutamides and - aspartamides as cholecystokinin antagonists

INVENTOR(S): Nadzan, Alex M.; Lin, Chun Wei; Kerwin, James F., Jr.
PATENT ASSIGNEE(S): Abbott Laboratories, USA

SOURCE: Eur. Pat. Appl., 66 pp.

CODRN: EPXXDM

DOCUMENT TYPE: Patent

LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
EP 308845	A1	19890329	EP 1988-115462	19880921 <<
R: ES, GR				
US 4971978	A	19901120	US 1988-234525	19880822 <<
WO 8902431	A1	19890323	WO 1988-US3181	19880921 <<
M: JP				
RM: BE, CH, DE, FR, GB, IT, NL, SE				
US 5128346	A	19920707	US 1980-571945	19800823 <<
PRIORITY APPLN. INFO.:			US 1987-9986	A 19870921
			US 1988-234525	A 19880822

OTHER SOURCE(S): MARPAT 111:134747

AB ArX1X2NR1 CH[(CH₂)nR4] CONR1R2 (I, R1 = H, Cl-alkyl, cycloalkyl, cyanoalkyl, adamantan, carbamoylalkyl, etc.; R2R2N = morpholino, pyrrolidiny, piperaziny, piperidino, etc.; R3 = H, alkyl, cycloalkyl, alkenyl, (substituted) arylalkyl, heterocyclylalkyl; R4 = tetrazolyl, acyl, Ar = heterocycl; XI = (CH₂)_n, OCH₂, SCH₂, NH = (substituted) alkynyl; X2 = CO, CS, SO₂; m = 0-4; n = 1-3), useful as cholecystokinin (CCK) antagonists, were prepared. H-Glu(OBz1)-N[(CH₂)_mE]2.HCl (preparation given) and N-methylmorpholine in DMF at 0° were treated successively with indole-2-carboxylic acid, 1-hydroxybenzotriazole, and 1-ethyl-3-(dimethylaminopropyl)carbodiimide. The mixture was allowed to warm to room temperature and stirred overnight and the product was debenzylated with Pd/C/cyclohexadiene to give N-(2-indolylcarbonyl)-L-glutamine di-N-pentylamide. It inhibit specific [¹²⁵I]-Bolton-Hunter CCK-8 pancreatic receptor binding with IC₅₀'s of 5.4-820 nm.

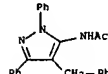
IT 122667-84-5P

RL: SPN (Synthetic preparation); PREP (Preparation)
 (preparation of an cholecystokinin antagonist)

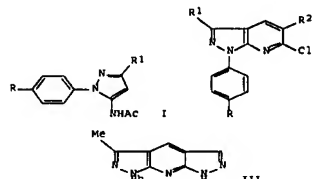
RN 122667-84-9 HCPLUS

CN Pentanoic acid, 5-((1,3-diphenyl-1H-pyrazol-5-yl)amino)-4-((1H-indol-2-ylcarbonyl)amino)-5-oxo-, (R)- (9CI) (CA INDEX NAME)

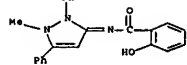
Absolute stereochemistry.



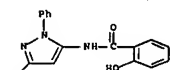
L5 ANSWER 46 OF 92 HCPLUS COPYRIGHT 2007 ACS on STN
ACCESSION NUMBER: 1982:217756 HCPLUS Full-text
DOCUMENT NUMBER: 96:217756
TITLE: Aminopyrazoles. II. Synthesis of pyrazolo[3,4,-b]pyridines via Vilsmeier-Haack reaction of 5-acetaminopyrazoles
AUTHOR(S): Simay, Antal; Takacs, Kalman; Toth, Laszlo
CORPORATE SOURCE: Res. Dep., Chinoin Pharm. Chem. Works, Budapest, Hung.
SOURCE: Acta Chimica Academiae Scientiarum Hungaricae (1982), 109(2), 175-87
CODEN: ACASAZ; ISSN: 0001-5407
DOCUMENT TYPE: Journal
LANGUAGE: English
OTHER SOURCE(S): CASREACT 96:217756
GI



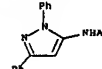
AB Vilsmeier-Haack reaction of pyrazoles I (R = H, Cl, NO₂, R₁ = H; R = H, R₁ = Me, Ph) gave 8-25% II (R₂ = H) and 35-53% III (R₂ = CHO). II (R = H, R₁ = Me, R₂ = CHO) was subjected to various reactions, including the formation of III.
IT 69730-07-0
RL: RCT (Reactant); RACT (Reactant or reagent)
(Vilsmeier-Haack reaction of)
RN 69730-07-0 HCPLUS
CN Acetamide, N-(1,3-diphenyl-1H-pyrazol-5-yl)- (9CI) (CA INDEX NAME)



IT 70803-13-3
RL: RCT (Reactant); RACT (Reactant or reagent)
(reaction of, with Me salicylate)
RN 70803-13-3 HCPLUS
CN Benzamide, N-(1,3-diphenyl-1H-pyrazol-5-yl)-2-hydroxy- (9CI) (CA INDEX NAME)



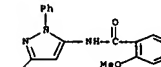
L5 ANSWER 48 OF 92 HCPLUS COPYRIGHT 2007 ACS on STN
ACCESSION NUMBER: 1981:538523 HCPLUS Full-text
DOCUMENT NUMBER: 95:138523
TITLE: Stability of packaged solid dosage forms. V. Prediction of the effect of aging on the disintegration of packaged tablets influenced by moisture and heat.
AUTHOR(S): Nakabayashi, Kiyoshi; Shimamoto, Tsugio; Mima, Hiroyuki; Okada, Jutaro
CORPORATE SOURCE: Cent. Res. Div., Takeda Chem. Ind., Ltd., Osaka, 532, Japan
SOURCE: Chemical & Pharmaceutical Bulletin (1981), 29(7), 2051-6
DOCUMENT TYPE: Journal
LANGUAGE: English
AB The effects of moisture and temperature on the disintegration time of a tablet containing gelatin as binder were investigated under accelerated conditions. The higher the ambient temperature and the moisture content of the tablet, the longer the disintegration time. Among several kinetic models investigated, a half-order reaction model was most suitable, where the ratio of the disintegration time of the aged samples to that of the initial ones was taken as a variable to be predicted. The effects of moisture and heat on the disintegration time ratio were analyzed by a multiple regression technique on the basis of the Carstensen's equation. In order to estimate the effect of aging on the disintegration time ratio, tablets in several kinds of moisture-semipermeable packages, including an overwrapped package, were examined in an artificial climate lab. The effect of aging could be predicted by an iterative calculation through a math. model in which the kinetics of the increase in the disintegration time ratio was combined with the moisture permeability of the packages. The simulated values could represent the observed data fairly well, although the confidence intervals of the predicted values were rather wide, owing to variances of the exptl. data obtained.
IT 69730-29-1
RL: BIOL (Biological study)
(tablet, disintegration of, moisture and storage temperature effect on)
RN 20170-20-1 HCPLUS



L5 ANSWER 47 OF 92 HCPLUS COPYRIGHT 2007 ACS on STN
ACCESSION NUMBER: 1981:569066 HCPLUS Full-text
DOCUMENT NUMBER: 95:169066
TITLE: Studies on the synthesis of heterocyclic compounds. Part VI. The action of methyl salicylate on some 5-aminopyrazoles
AUTHOR(S): Daidone, Giuseppe; Plescia, Salvatore
CORPORATE SOURCE: Ist. Chim. Farm. Tossicolog., Univ. Palermo, Palermo, 32-90123, Italy
SOURCE: Journal of Heterocyclic Chemistry (1981), 18(4), 747-50
DOCUMENT TYPE: Journal
LANGUAGE: English
OTHER SOURCE(S): CASREACT 95:169066
GI

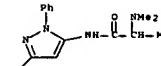


AB Refluxing 5-aminopyrazoles I (R = H, Me, Ph; R₁ = H) with o-HOC₆H₄CO₂Me gave I (R₁ = O-MeOCH₂CO₂Me) and II.
IT 79442-81-2P 79442-84-5P
RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)
(preparation and hydrolysis of)
RN 79442-81-2 HCPLUS
CN Benzamide, N-(1,3-diphenyl-1H-pyrazol-5-yl)-2-methoxy- (9CI) (CA INDEX NAME)

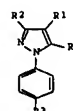


RN 79442-84-5 HCPLUS
CN Benzamide, N-(1,3-diphenyl-1H-pyrazol-5-yl)-2-methoxy- (9CI) (CA INDEX NAME)

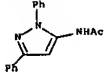
CN Propanamide, 2-(dimethylamino)-N-(1,3-diphenyl-1H-pyrazol-5-yl)- (9CI) (CA INDEX NAME)



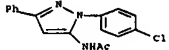
L5 ANSWER 49 OF 92 HCPLUS COPYRIGHT 2007 ACS on STN
ACCESSION NUMBER: 1981:442979 HCPLUS Full-text
DOCUMENT NUMBER: 95:42979
TITLE: Vilsmeier-Haack reaction of 5-amino- and 5-acetylaminopyrazoles
AUTHOR(S): Simay, A.; Takacs, K.; Horvath, K.; Dvortsak, P.
CORPORATE SOURCE: Res. Dep., Chinoin Pharm. Chem. Works, Budapest, Hung.
SOURCE: Acta Chimica Academiae Scientiarum Hungaricae (1980), 105(2), 127-39
CODEN: ACASAZ; ISSN: 0001-5407
DOCUMENT TYPE: Journal
LANGUAGE: English
OTHER SOURCE(S): CASREACT 95:42979
GI



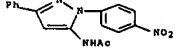
AB Vilsmeier-Haack reaction of aminopyrazoles I (R = NH₂, R₁ = H, R₂ = H, Me, Ph, R₃ = H, Cl, NO₂, Me, OMe) gave I (R = N:CHNR₄2, R₁ = CHO, NR₄2 = NMe₂, piperidino). The intermediates I (R = N:CHNR₄2, R₁ = H, CH₂N+R₄X-, X = Cl, PO₂C₁₂, ClO₄) were isolated. I (R = NHCHO, R₁ = H) similarly gave I (R = N:CHNR₄2, R₁ = CHO, H). I (R = N:CHNR₄2, R₁ = CHO) were converted into I (R₁ = CH:NPh, CH:NNH₂, CH:NNHC₁₂NH₂) and I [R₁ = N:CH:NCH, N:CH(N₂O):CH].
IT 69730-07-0P 77745-65-1P 77745-96-8P
RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)
(preparation and Vilsmeier-Haack reaction of)
RN 69730-07-0 HCPLUS
CN Acetamide, N-(1,3-diphenyl-1H-pyrazol-5-yl)- (9CI) (CA INDEX NAME)



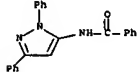
RN 77746-85-1 HCAPLUS
CN Acetamide, N-[1-(4-chlorophenyl)-3-phenyl-1H-pyrazol-5-yl]- (9CI) (CA INDEX NAME)



RN 77746-87-3 HCAPLUS
CN Acetamide, N-[1-(4-nitrophenyl)-3-phenyl-1H-pyrazol-5-yl]- (9CI) (CA INDEX NAME)

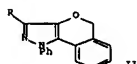


RN 77746-90-8 HCAPLUS
CN Benzamide, N-(1,3-diphenyl-1H-pyrazol-5-yl)- (9CI) (CA INDEX NAME)



L5 ANSWER 50 OF 92 HCAPLUS COPYRIGHT 2007 ACS on STN
ACCESSION NUMBER: 1981:212395 HCAPLUS Full-text
DOCUMENT NUMBER: 94:212395
TITLE: Studies on the synthesis of heterocyclic compounds.
Part IV. Further investigation of the Pechmann reaction with some pyrazole derivatives
AUTHOR(S): Daidone, Giuseppe; Plescia, Salvatore; Fabra, Jole
COPARTNERS: Ist. Chim. Farm. Tossicol., Univ. Via Archirafi,
Palermo, 32-90123, Italy
SOURCE: Journal of Heterocyclic Chemistry (1980),
17(7), 1409-11
CODEN: JHTCAD; ISSN: 0022-152X
DOCUMENT TYPE: Journal

LANGUAGE: English
OTHER SOURCE(S): CASREACT 94:121395
GI



AB Thermal decomposition of the diazonium sulfate derived from N-methyl-(1-phenyl-3-methylpyrazol-5-yl)-2-aminobenzamide afforded products formulated as 1-phenyl-3-methyl[2]benzopyran-4,3-cipyrazol-5-one (yield 10%), 1,4-dimethyl-3-phenylpyrazolo[3,4-c]isoquinolin-5-one (yield 10%), N-methyl-(1-phenyl-3-methylpyrazol-5-yl)-2-hydroxybenzamide (yield 8%) and 4'-hydroxy-2,3'-dimethyl-1'-phenylspiro[isindoline-1,5'-[2]pyrazolin]-3-one (I) (yield 20%). Decomposition of the diazonium sulfate derived from N-methyl-(1,3-diphenylpyrazol-5-yl)-2-aminobenzamide gave products formulated as 7,9-dimethyldibenzo[e,g]pyrazolo[1,5-a][1,3]diazocin-10(9H)-one (yield 8%), 4-methyl-1,3-diphenylpyrazolo[3,4-c]isoquinolin-5-one (yield 7%) and 4'-hydroxy-2-methyl-1',3'-diphenylspiro[isindoline-1,5'-[2]pyrazolin]-3-one (II) (yield 10%). The spiro compds. I and II underwent thermal and acid-catalyzed conversion into the hitherto unknown 2-benzopyran-4,3-cipyrazole ring system III (R = Me, Ph) in good yield.

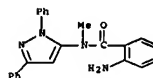
IT 69730-14-9

RL: RCT (Reactant); RACT (Reactant or reagent)

(preparation of diazonium sulfate from, decomposition of)

RN 69730-14-9 HCAPLUS

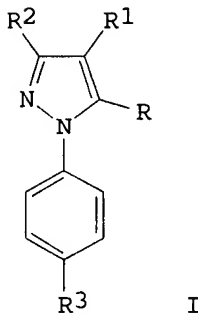
CN Benzamide, 2-amino-N-(1,3-diphenyl-1H-pyrazol-5-yl)-N-methyl- (9CI) (CA INDEX NAME)



> log hold		SINCE FILE	TOTAL
COST IN U.S. DOLLARS		ENTRY	SESSION
FULL ESTIMATED COST		341.50	514.26
DISCOUNT AMOUNTS (FOR QUALIFYING ACCOUNTS)		SINCE FILE	TOTAL
CA SUBSCRIBER PRICE		39.00	39.00

SESSION WILL BE HELD FOR 120 MINUTES
STN INTERNATIONAL SESSION SUSPENDED AT 09:43:53 ON 23 MAY 2007

L5 ANSWER 49 OF 92 HCAPLUS COPYRIGHT 2007 ACS on STN
 ACCESSION NUMBER: 1981:442979 HCAPLUS <<LOGINID::20070523>>
 DOCUMENT NUMBER: 95:42979
 TITLE: Vilsmeier-Haack reaction of 5-amino- and
 5-acylaminopyrazoles
 AUTHOR(S): Simay, A.; Takacs, K.; Horvath, K.; Dvortsak, P.
 CORPORATE SOURCE: Res. Dep., Chinoin Pharm. Chem. Works, Budapest, Hung.
 SOURCE: Acta Chimica Academiae Scientiarum Hungaricae (1980), 105(2), 127-39
 DOCUMENT TYPE: CODEN: ACASA2; ISSN: 0001-5407
 LANGUAGE: English
 OTHER SOURCE(S): CASREACT 95:42979
 GI



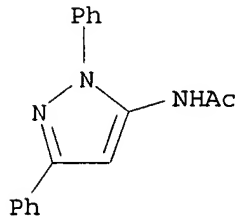
AB Vilsmeier-Haack reaction of aminopyrazoles I ($R = \text{NH}_2$, $R1 = \text{H}$, $R2 = \text{H}$, Me , Ph , $R3 = \text{H}$, Cl , NO_2 , Me , OMe) gave I ($R = \text{N:CHNR42}$, $R1 = \text{CHO}$, $\text{NR42} = \text{NMe}_2$, piperidino). The intermediates I ($R = \text{N:CHNR42}$, $R1 = \text{H}$, CH:N+R42X- , $X = \text{Cl}$, PO_2Cl_2 , ClO_4) were isolated. I ($R = \text{NHCHO}$, NHAc , NHBz , $R1 = \text{H}$) similarly gave I ($R = \text{N:CHNR42}$, $R1 = \text{CHO}$, H). I ($R = \text{N:CHNMe}_2$, $R1 = \text{CHO}$) were converted into I ($R1 = \text{CH:NPh}$, CH:NNHPh , CH:NNHCONH_2 , CH:NNHCSNH_2) and I [$\text{RR1} = \text{N:CHN:CH}$, N:CHN(O):CH].

IT 69730-07-0P 77746-85-1P 77746-87-3P
 77746-90-8P
 RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(preparation and Vilsmeier-Haack reaction of)

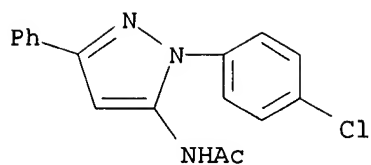
RN 69730-07-0 HCAPLUS

CN Acetamide, N-(1,3-diphenyl-1H-pyrazol-5-yl)- (9CI) (CA INDEX NAME)



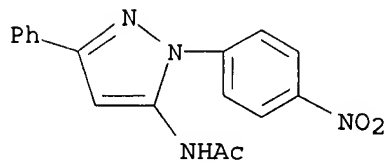
RN 77746-85-1 HCAPLUS

CN Acetamide, N-[1-(4-chlorophenyl)-3-phenyl-1H-pyrazol-5-yl]- (9CI) (CA INDEX NAME)



RN 77746-87-3 HCAPLUS

CN Acetamide, N-[1-(4-nitrophenyl)-3-phenyl-1H-pyrazol-5-yl]- (9CI) (CA INDEX NAME)



RN 77746-90-8 HCAPLUS

CN Benzamide, N-(1,3-diphenyl-1H-pyrazol-5-yl)- (9CI) (CA INDEX NAME)

